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(57) Abstract

Novel chimeric nucleic acids, encoding chimeric Borrelia proteins consisting of at least two antigenic polypeptides from corresponding and/or non-corresponding proteins from the same and/or different species of Borrelia, are disclosed. Chimeric proteins encoded by the nucleic acid sequences are also disclosed. The chimeric proteins are useful as vaccine immunogens against Lyme borreliosis, as well as for immunodiagnostic reagents.

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CHIMERIC PROTEINS COMPRISING BORRELIA POLYPEPTIDES; USES THEREFOR

Background of the Invention

Lyme borreliosis is the most common tick-borne 5 infectious disease in North America, Europe, and northern Asia. The causative bacterial agent of this disease, Borrelia burgdorferi, was first isolated and cultivated in 1982 (Burgdorferi, W.A. et al., Science 216: 1317-1319 (1982); Steere, A.R. et al., N. Engl. J. 10 Med. 308: 733-740 (1983)). With that discovery, a wide array of clinical syndromes, described in both the European and American literature since the early 20th century, could be attributed to infection by B. burgdorferi (Afzelius, A., Acta Derm. Venereol. 2: 120-15 125 (1921); Bannwarth, A., Arch. Psychiatr. Nervenkrankh. 117: 161-185 (1944); Garin, C. and A. Bujadouz, J. Med. Lyon 71: 765-767 (1922); Herxheimer, K. and K. Hartmann, Arch. Dermatol. Syphilol. 61: 57-76, 255-300 (1902)).

The immune response to B. burgdorferi is characterized by an early, prominent, and persistent humoral response to the end of lagellar protein, p41 (fla), and to a protein constituent of the protoplasmic cylinder, p93 (Szczepanski, A., and J.L. Benach,

Microbiol. Rev. 55:21 (1991)). The p41 flagellin antigen is an immunodominant protein; however, it shares significant homology with flagellins of other microorganisms and therefore is highly cross reactive. The p93 antigen is the largest immunodominant antigen of

30 B. burgdorferi. Both the p41 and p93 proteins are physically cryptic antigens, sheathed from the immune system by an outer membrane whose major protein constituents are the outer surface proteins A and B

(OspA and OspB). OspA is a basic lipoprotein of approximately 31 kd, which is encoded on a large linear plasmid along with OspB, a basic lipoprotein of approximately 34 kd (Szczepanski, A., and J.L. Benach, 5 Microbiol. Rev. 55:21 (1991)). Analysis of isolates of B. burgdorferi obtained from North America and Europe has demonstrated that OspA has antigenic variability, and that several distinct groups can be serologically and genotypically defined (Wilske, B., et al., World J. 10 Microbiol. 7: 130 (1991)). Other Borrelia proteins demonstrate similar antigenic variability. Surprisingly, the immune response to these outer surface proteins tends to occur late in the disease, if at all (Craft, J. E. et al., <u>J. Clin Invest. 78</u>: 934-939 15 (1986); Dattwyler, R.J. and B.J. Luft, Rheum. Clin. North Am. 15: 727-734 (1989)). Furthermore, patients acutely and chronically infected with B. burgdorferi respond variably to the different antigens, including OspA, OspB, OspC, OspD, p39, p41 and p93.

Vaccines against Lyme borreliosis have been attempted. Mice immunized with a recombinant form of OspA are protected from challenge with the same strain of B. burgdorferi from which the protein was obtained (Fikrig, E., et al., Science 250: 553-556 (1990)).

25 Furthermore, passively transferred anti-OspA monoclonal antibodies (Mabs) have been shown to be protective in mice, and vaccination with a recombinant protein induced protective immunity against subsequent infection with the homologous strain of B.burgdorferi (Simon, M.M., et al., J. Infect. Dis. 164: 123 (1991)). Unfortunately, immunization with a protein from one strain does not necessarily confer resistance to a heterologous strain (Fikrig, E. et al., J. Immunol. 7: 2256-1160 (1992)), but rather, is limited to the homologous 'species' from

35 which the protein was prepared. Furthermore,

immunization with a single protein from a particular strain of Borrelia will not confer resistance to that strain in all individuals. There is considerable variation displayed in OspA and OspB, as well as p93, including the regions conferring antigenicity. Therefore, the degree and frequency of protection from vaccination with a protein from a single strain depend upon the response of the immune system to the particular variation, as well as the frequency of genetic variation in B. burgdorferi. Currently, a need exists for a vaccine which provides immunogenicity across species and to more epitopes within a species, as well as immunogenicity against more than one protein.

Summary of the Invention

The current invention pertains to chimeric Borrelia 15 proteins which include two or more antigenic Borrelia polypeptides which do not occur naturally (in nature) in the same protein in Borrelia, as well as the nucleic acids encoding such chimeric proteins. The antigenic 20 polypeptides incorporated in the chimeric proteins are derived from any Borrelia protein from any strain of Borrelia, and include outer surface protein (Osp) A, OspB, OspC, OspD, p12, p39, p41, p66, and p93. The proteins from which the antigenic polypeptides are derived can be from the same strain of Borrelia, from 25 different strains, or from combinations of proteins from the same and from different strains. If the proteins from which the antigenic polypeptides are derived are OspA or OspB, the antigenic polypeptides can be derived 30 from either the portion of the OspA or OspB protein present between the amino terminus and the conserved tryptophan of the protein (referred to as a proximal portion), or the portion of the OspA or OspB protein present between the conserved tryptophan of the protein

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and the carboxy terminus (referred to as a distal portion). Particular chimeric proteins, and the nucleotide sequences encoding them, are set forth in Figures 23-37 and 43-46.

The chimeric proteins of the current invention provide antigenic polypeptides of a variety of Borrelia strains and/or proteins within a single protein. proteins are particularly useful in immunodiagostic assays to detect the presence of antibodies to native 10 Borrelia in potentially infected individuals as well as to measure T-cell reactivity, and can therefore be used as immunodiagnostic reagents. The chimeric proteins of the current invention are additionally useful as vaccine immunogens against Borrelia infection.

For a better understanding of the present invention together with other and further objects, reference is made to the following description, taken together with the accompanying drawings.

Brief Description of the Drawings

20 Figure 1 summarizes peptides and antigenic domains localized by proteolytic and chemical fragmentation of OspA.

Figure 2 is a comparison of the antigenic domains depicted in Figure 1, for OspA in nine strains of B. 25 burgdorferi.

Figure 3 is a graph depicting a plot of weighted polymorphism versus amino acid position among 14 OspA variants. The marked peaks are: a) amino acids 132-145; b) amino acids 163-177; c) amino acids 208-221. 30 lower dotted line at polymorphism value 1.395 demarcates statistically significant excesses of polymorphism at p The upper dotted line at 1.520 is the same, except that the first 29 amino acids at the monomorphic N-terminus have been removed from the original analysis.

Figure 4 depicts the amino acid alignment of residues 200 through 220 for OspAs from strains B31 and K48 as well as for the site-directed mutants 613, 625, 640, 613/625, and 613/640. Arrow indicates Trp216.

5 Amino acid changes are underlined.

Figure 5 is a helical wheel projection of residues 204-217 cf B31 OspA. Capital letters indicate hydrophobic residues; lower case letters indicate hydrophilic residues; +/- indicate positively/negatively charged residues. Dashed line indicates division of the alpha-helix into hydrophobic arc (above the line) and polar arc (below the line). Adapted from France et al. (Biochem. Biophys. Acta 1120: 59 (1992)).

Figure 6 depicts a phylogenic tree for strains of

Borrelia described in Table I. The strains are as
follows: 1 = B31; 2 = Pka1; 3 = ZS7; 4 = N40; 5 =
25015; 6 = K48; 7 = DK29; 8 = PHei; 9 = Ip90; 10 =
PTrob; 11 = ACAI; 12 = PGau; 13 = Ip3; 14 = PBo; 15 =
PKo.

Figure 7 depicts the nucleic acid sequence of OspA-B31 (SEW ID NO. 6), and the encoded protein sequence (SEQ ID NO. 7).

Figure 8 depicts the nucleic acid sequence of OspA-K48 (SEQ ID NO. 8), and the encoded protein sequence (SEQ ID NO. 9).

Figure 9 depicts the nucleic acid sequence of OspA-PGau (SEQ ID NO. 10), and the encoded protein sequence (SEQ ID NO. 11).

Figure 10 depicts the nucleic acid sequence of OSPA-25015 (SEQ ID NO. 12), and the encoded protein sequence (SEQ ID NO. 13).

Figure 11 depicts the nucleic acid sequence of OspB-B31 (SEQ ID NO. 21), and the encoded protein sequence (SEQ ID NO. 22).

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Figure 12 depicts the nucleic acid sequence of OspC-B31 (SEQ ID NO. 29), and the encoded protein sequence (SEQ ID NO. 30).

Figure 13 depicts the nucleic acid sequence of OspC-K48 (SEQ ID NO. 31), and the encoded protein sequence (SEQ ID NO. 32).

Figure 14 depicts the nucleic acid sequence of OspC-PKo (SEQ ID NO. 33), and the encoded protein sequence (SEQ ID NO. 34).

Figure 15 depicts the nucleic acid sequence of OspC-pTrob (SEQ ID NO. 35) and the encoded protein sequence (SEQ ID NO. 36).

Figure 16 depicts the nucleic acid sequence of p93-B31 (SEQ ID NO. 65) and the encoded protein sequence (SEQ ID NO. 66).

Figure 17 depicts the nucleic acid sequence of p93-K48 (SEQ ID NO. 67).

Figure 18 depicts the nucleic acid sequence of p93-PBo (SEQ ID NO. 69).

Figure 19 depicts the nucleic acid sequence of p93pTrob (SEQ ID NO. 71).

Figure 20 depicts the nucleic acid sequence of p93-pGau (SEQ ID NO. 73).

Figure 21 depicts the nucleic acid sequence of p93- 25-25015 (SEQ ID NO. 75).

Figure 22 depicts the nucleic acid sequence of p93-pKo (SEQ ID NO. 77).

Figure 23 depicts the nucleic acid sequence of the OspA-K48/OspA-PGau chimer (SEQ ID NO. 85) and the encoded chimeric protein sequence (SEQ ID No. 86).

Figure 24 depicts the nucleic acid sequence of the OspA-B31/OspA-PGau chimer (SEQ ID NO. 88) and the encoded chimeric protein sequence (SEQ ID NO. 89).

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Figure 25 depicts the nucleic acid sequence of the OspA-B31/OspA-K48 chimer (SEQ ID NO. 91) and the encoded chimeric protein sequence (SEQ ID NO. 92).

Figure 26 depicts the nucleic acid sequence of the OspA-B31/OspA-25015 chimer (SEQ ID NO. 94) and the encoded chimeric protein sequence (SEQ ID NO. 95).

Figure 27 depicts the nucleic acid sequence of the OspA-K48/OspA-B31/OspA-K48 chimer (SEQ ID NO. 97) and the encoded chimeric protein sequence (SEQ ID NO. 98).

Figure 28 depicts the nucleic acid sequence of the OspA-B31/OspA-K48/OspA-B31/OspA-K48 chimer (SEQ ID NO. 100) and the encoded chimeric protein sequence (SEQ ID NO. 101).

Figure 29 depicts the nucleic acid sequence of the OspA-B31/OspB-B31 chimer (SEQ ID NO. 103) and the encoded chimeric protein sequence (SEQ ID NO. 104).

Figure 30 depicts the nucleic acid sequence of the OspA-B31/OspB-B31/OspC-B31 chimer (SEQ ID NO. 106) and the encoded chimeric protein sequence (SEQ ID NO. 107).

20 Figure 31 depicts the nucleic acid sequence of the OspC-B31/OspA-B31/OspB-B31 chimer (SEQ ID NO. 109) and the encoded chimeric protein sequence (SEQ ID NO. 110).

Figure 32 depicts the nucleic acid sequence of the OspA-B31/p93-B31 chimer (SEQ ID NO. 111) and the encoded chimeric protein sequence (SEQ ID NO. 112).

Figure 33 depicts the nucleic acid sequence of the OspB-B31/p41-B31 (122-234) chimer (SEQ ID NO. 113) and the encoded chimeric protein sequence (SEQ ID NO. 114).

Figure 34 depicts the nucleic acid sequence of the OspB-B31/p41-B31 (122-295) chimer (SEQ ID NO. 115) and the encoded chimeric protein sequence (SEQ ID NO. 116).

Figure 35 depicts the nucleic acid sequence of the OspB-B31/p41-B31 (140-234) chimer (SEQ ID NO. 117) and the encoded chimeric protein sequence (SEQ ID NO. 118).

Figure 36 depicts the nucleic acid sequence of the OspB-B31/p41-B31 (140-295) chimer (SEQ ID NO. 119) and the encoded chimeric protein sequence (SEQ ID NO. 120).

Figure 37 depicts the nucleic acid sequence of the 5 OspB-B31/p41-B31 (122-234)/OspC-B31 chimer (SEQ ID NO. 121) and the encoded chimeric protein sequence (SEQ ID NO. 122).

Figure 38 depicts an alignment of the nucleic acid sequences for OspC-B31 (SEQ ID NO. 29), OspC-PKo (SEQ ID NO. 33), OspC-pTrob (SEQ ID NO. 35), and OspC-K48 (SEQ ID NO. 31). Nucleic acids which are identical to those in the lead nucleic acid sequence (here, OspC-B31) are represented by a period (.); differing nucleic acids are shown in lower case letters.

Figure 39 depicts an alignment of the nucleic acid sequences for OspD-pBO (SEQ ID NO. 123), OspD-PGau (SEQ ID NO. 124), OspD-DK29 (SEQ ID NO. 125), and OspD-K48 (SEQ ID NO. 126). Nucleic acids which are identical to those in the lead nucleic acid sequence (here, OspD-pBo) are represented by a period (.); differing nucleic acids are shown in lower case letters.

Figure 40 depicts the nucleic acid sequence of p41-B31 (SEq ID NO. 127) and then encoded protein sequence (SEQ ID NO. 128).

Figure 41 depicts an alignment of the nucleic acid sequences for p41-B31 (SEQ ID NO. 127), p41-pKa1 (SEQ ID NO. 129), p41-PGau (SEQ ID NO. 51), p41-PBo (SEQ ID NO. 130), p41-DK29 (SEQ ID NO. 53), and p41-PKo (SEQ ID NO. 131). Nucleic acids which are identical to those in the lead nucleic acid sequence (here, p41-B31) are represented by a period (.); differing nucleic acids are shown in lower case letters.

Figure 42 depicts an alignment of the nucleic acid sequences for OspA-B31 (SEQ ID NO. 6), OspA-pKa1 (SEQ ID NO. 132), OspA-N40 (SEQ ID NO. 133), OspA-ZS7 (SEQ ID

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NO. 134), OspA-25015 (SEQ ID NO. 12), OspA-pTrob (SEQ ID NO. 135), OspA-K48 (SEQ ID NO. 8), OspA-Hei (SEQ ID NO. 136), OspA-DK29 (SEQ ID NO. 49), OSpA-Ip90 (SEQ ID NO. 50), OspA-pBo (Seq ID NO. 55), OspA-Ip3 (SEQ ID NO. 56), OspA-PKo (SEQ ID NO. 57), OspA-ACAI (SEQ ID NO. 58), and OspA-PGau (SEQ ID NO. 10). Nucleic acids which are identical to those in the lead nucleic acid sequence (here, OspA-B31) are represented by a period (.); differing nucleic acids are shown in lower case letters.

Figure 43 depicts the nucleic acid sequence of the OspA-Tro/OspA-Bo chimer (SEQ ID NO. 137) and the encoded chimeric protein sequence (SEQ ID NO. 138).

Figure 44 depicts the nucleic acid sequence of the OspA-PGau/OspA-Bo chimer (SEQ ID NO. 139) and the encoded chimeric protein sequence (SEQ ID NO. 140).

Figure 45 depicts the nucleic acid sequence of the OspA-B31/OspA-PGau/OspA-B31/OspA-K48 chimer (SEQ ID NO. 141) and the encoded chimeric protein sequence (SEQ ID NO. 142).

Figure 46 depicts the nucleic acid sequence of the OspA-PGau/OspA-B31/OspA-K48 chimer (SEQ ID NO. 143) and the encoded chimeric protein sequence (SEQ ID NO. 144).

Detailed Description of the Invention

The current invention pertains to chimeric proteins comprising antigenic Borrelia polypeptides which do not occur in nature in the same Borrelia protein. The chimeric proteins are a combination of two or more antigenic polypeptides derived from Borrelia proteins. The antigenic polypeptides can be derived from different proteins from the same species of Borrelia, or different proteins from different Borrelia species, as well as from corresponding proteins from different species. As used herein, the term "chimeric protein" describes a protein comprising two or more polypeptides which are

derived from corresponding and/or non-corresponding native Borrelia protein. A polypeptide "derived from" a native Borrelia protein is a polypeptide which has an amino acid sequence the same as an amino acid sequence 5 present in a Borrelia protein, an amino acid sequence equivalent to the amino acid sequence of a naturally occurring Borrelia protein, or an amino acid sequence substantially similar to the amino acid sequence of a naturally occurring Borrelia protein (e.g., differing by 10 few amino acids) such as when a nucleic acid encoding a protein is subjected to site-directed mutagenesis. "Corresponding" proteins are equivalent proteins from different species or strains of Borrelia, such as outer surface protein A (OspA) from strain B31 and OspA from 15 strain K48. The invention additionally pertains to nucleic acids encoding these chimeric proteins.

As described below, Applicants have identified two separate antigenic domains of OspA and OspB which flank the sole conserved tryptophan present in OspA and in 20 OspB. These domains share cross-reactivity with different genospecies of Borrelia. The precise amino acids responsible for antigenic variability were determined through site-directed mutagenesis, so that proteins with specific amino acid substitutions are available for the development of chimeric proteins. Furthermore, Applicants have identified immunologically important hypervariable domains in OspA proteins, as described below in Example 2. The first hypervariable domain of interest for chimeric proteins, Domain A, includes amino acid residues 120-140 of OspA, the second hypervariable domain, Domain B, includes residues 150-180 and the third hypervariable domain, Domain C, includes residues 200-216 or 217 (depending on the position of the sole conserved tryptophan residue in the 35 OspA of that particular species of Borrelia) (see Figure 3). In addition, Applicants have sequenced the genes for several Borrelia proteins.

These discoveries have aided in the development of novel recombinant Borrelia proteins which include two or more amino acid regions or sequences which do not occur in the same Borrelia protein in nature. The recombinant proteins comprise polypeptides from a variety of Borrelia proteins, including, but not limited to, OspA, OspB, OspC, OspD, p12, p39, p41, p66, and p93.

10 Antigenically relevant polypeptides from each of a number of proteins are combined into a single chimeric protein.

In one embodiment of the current invention, chimers are now available which include antigenic polypeptides 15 flanking a tryptophan residue. The antigenic polypeptides are derived from either the proximal portion from the tryptophan (the portion of the OspA or OspB protein present between the amino terminus and the conserved tryptophan of the protein), or the distal 20 portion from the tryptophan (the portion of the OspA or OspB protein present between the conserved tryptophan of the protein and the carboxy terminus) in OspA and/or OspB. The resultant chimers can be OspA-OspA chimers (i.e., chimers incorporating polypeptides derived from 25 OspA from different strains of Borrelia), OspA-OspB chimers, or OspB-OspB chimers, and are constructed such that amino acid residues amino-proximal to an invariant tryptophan are from one protein and residues carboxyproximal to the invariant tryptophan are from the other 30 protein. For example, one available chimer consists of a polypeptide derived from the amino-proximal region of OspA from strain B31, followed by the tryptophan residue, followed by a polypeptide derived from the carboxy-proximal region of OspA from strain K48 (SEQ ID 35 NO. 92). Another available chimer includes a

polypeptide derived from the amino-proximal region of OspA from strain B31, and a polypeptide derived from the carboxy-proximal region of OspB from strain B31 (SEQ ID NO. 104). If the polypeptide proximal to the tryptophan of these chimeric proteins is derived from OspA, the proximal polypeptide can be further subdivided into the three hypervariable domains (Domains A, B, and C), each of which can be derived from OspA from a different strain of Borrelia. These chimeric proteins can further comprise antigenic polypeptides from another protein, in addition to the antigenic polypeptides flanking the tryptophan residue.

In another embodiment of the current invention, chimeric proteins are available which incorporate antigenic domains of two or more *Borrelia* proteins, such as Osp proteins (Osp A, B, C and/or D) as well as p12, p39, p41, p66, and/or p93.

The chimers described herein can be produced so that they are highly soluble, hyper-produced in E. coli, and non-lipidated. In addition, the chimeric proteins can be designed to end in an affinity tag (His-tag) to facilitate purification. The recombinant proteins described herein have been constructed to maintain high levels of antigenicity. In addition, recombinant proteins specific for the various genospecies of Borrelia that cause Lyme disease are now available, because the genes from each of the major genospecies have been sequenced; the sequences are set forth below. These recombinant proteins with their novel biophysical and antigenic properties will be important diagnostic reagent and vaccine candidates.

The chimeric proteins of the current invention are advantageous in that they retain specific reactivity to monoclonal and polyclonal antibodies against wild-type Borrelia proteins, are immunogenic, and inhibit the

growth or induce lysis of Borrelia in vitro.

Furthermore, in some embodiments, the proteins provide antigenic domains of two or more Borrelia strains and/or proteins within a single protein. Such proteins are particularly useful in immuno-diagostic assays. For example, proteins of the present invention can be used as reagents in assays to detect the presence of antibodies to native Borrelia in potentially infected individuals. These proteins can also be used as immunodiagnostic reagents, such as in dot blots, Western blots, enzyme linked immunosorbed assays, or agglutination assays. The chimeric proteins of the present invention can be produced by known techniques, such as by recombinant methodology, polymerase chain reaction, or mutagenesis.

Furthermore, the proteins of the current invention are useful as vaccine immunogens against Borrelia infection. Because Borrelia has been shown to be clonal, a protein comprising antigenic polypeptides from 20 a variety of Borrelia proteins and/or species, will provide immunoprotection for a considerable time when used in a vaccine. The lack of significant intragenic recombination, a process which might rapidly generate novel epitopes with changed antigenic properties, 25 ensures that Borrelia can only change antigenic type by accumulating mutational change, which is slow when compared with recombination in generating different antigenic types. The chimeric protein can be combined with a physiologically acceptable carrier and 30 administered to a vertebrate animal through standard methods (e.g., intravenously or intramuscularly, for example).

The current invention is illustrated by the following Examples, which are not to be construed to be limiting in any way.

Example 1. Purification of Borrelia burgorferi Outer Surface Protein A and Analysis of Antibody Binding Domains

This example details a method for the purification of large amounts of native outer surface protein A (OspA) to homogeneity, and describes mapping of the antigenic specificities of several anti-OspA MAbs. OspA was purified to homogeneity by exploiting its resistance to trypsin digestion. Intrinsic labeling with ¹⁴C-palmitic acid confirmed that OspA was lipidated, and partial digestion established lipidation at the aminoterminal cysteine of the molecule.

The reactivity of seven anti-OspA murine monoclonal antibodies to nine different Borrelia isolates was ascertained by Western blot analysis. Purified OspA was fragmented by enzymatic or chemical cleavage, and the monoclonal antibodies were able to define four distinct immunogenic domains (see Figure 1). Domain 3, which included residues 190-220 of OspA, was reactive with protective antibodies known to agglutinate the organism in vitro, and included distinct specificities, some of which were not restricted to a genotype of B. burgdorferi.

A. Purification of Native OspA

Detergent solubilization of B. burgdorferi strips the outer surface proteins and yields partially-purified preparations containing both OspA and outer surface 5 protein B (Osp B) (Barbour, A.G. et al., Infect. Immun. 52 (5): 549-554 (1986); Coleman, J.L. and J.L. Benach, J Infect. Dis. 155 (4): 756-765 (1987); Cunningham, T.M. et al., Ann. NY Acad. Sci. 539: 376-378 (1988); Brandt, M.E. et al., <u>Infect. Immun. 58</u>: 983-991 (1990); Sambri, 10 V. and R. Cevenini, Microbiol. 14:307-314 (1991)). Although both OspA and OspB are sensitive to proteinase K digestion, in contrast to OspB, OspA is resistant to cleavage by trypsin (Dunn, J. et al., Prot. Exp. Purif. 1: 159-168 (1990); Barbour, A.G. et al., <u>Infect. Immun.</u> 15 <u>45</u>:94-100 (1984)). The relative insensitivity to trypsin is surprising in view of the fact that Osp A has a high (16% for B31) lysine content, and may relate to the relative configuration of Osp A and B in the outer membrane.

Intrinsic Radiolabeling of Borrelia
Labeling for lipoproteins was performed as
described by Brandt et al. (<u>Infect. Immun. 58</u>:983-991
(1990)). ¹⁴C-palmitic acid (ICN, Irvine, California) was
added to the BSK II media to a final concentration of
0.5 μCi per milliliter (ml). Organisms were cultured at
34°C in this medium until a density of 10⁸ cells per ml
was achieved.

Purification of OspA Protein from Borrelia Strain B31

Borrelia burgdorferi, either ¹⁴C-palmitic acid
labeled or unlabeled, were harvested and washed as
described (Brandt, M.E. et al., <u>Infect. Immun. 58</u>:983991 (1990)). Whole organisms were trypsinized according

to the protocol of Barbour et al. (Infect. Immun. 45:94-100 (1984)) with some modifications. The pellet was suspended in phosphate buffered saline (PBS, 10mM, pH 7.2), containing 0.8% tosyl-L-phenylalanine chloromethyl 5 ketone (TPCK)-treated trypsin (Sigma, St. Louis, Missouri), the latter at a ratio of 1 μ g per 10 8 cells. Reaction was carried out at 25°C for 1 hour, following which the cells were centrifuged. The pellet was washed in PBS with 100 μ g/ml phenylmethylsulfonyl fluoride 10 (PMSF). Triton X-114 partitioning of the pellet was carried out as described by Brandt et al. (Infect. Immun. 58:983-991 (1990)). Following trypsin treatment, cells were resuspended in ice-cold 2% (V/V) Triton X-114 in PBS at 109 cells per ml. The suspension was rotated 15 overnight at 4°C, and the insoluble fraction removed as a pellet after centrifugation at 10,000 X g for 15 minutes at 4°C. The supernatant (soluble fraction) was incubated at 37°C for 15 minutes and centrifuged at room temperature at 1000 X g for 15 minutes to separate the 20 aqueous and detergent phases. The aqueous phase was decanted, and ice cold PBS added to the lower Triton phase, mixed, warmed to 37°C, and again centrifuged at 1000 X g for 15 minutes. Washing was repeated twice Finally, detergent was removed from the 25 preparation using a spin column of Bio-beads SM2 (BioRad, Melville, New York) as described (Holloway, P.W., Anal. Biochem. 53:304-308 (1973)).

Ion exchange chromatography was carried out as described by Dunn et al. (Prot. Exp. Purif. 1: 159-168 (1990)) with minor modifications. Crude OspA was dissolved in buffer A (1% Triton X-100, 10mM phosphate buffer (pH 5.0)) and loaded onto a SP Sepharose resin (Pharmacia, Piscataway, New Jersey), pre-equilibrated with buffer A at 25°C. After washing the column with 10

bed-volumes of buffer A, the bound OspA was eluted with buffer B (1% Triton X-100, 10mM phosphate buffer (pH 8.0)). OspA fractions were detected by protein assay using the BCA method (Pierce, Rockford, Illinois), or as radioactivity when intrinsically labeled material was fractionated. Triton X-100 was removed using a spin column of Bio-beads SM2.

This method purifies OspA from an outer surface membrane preparation. In the absence of trypsin
10 treatment, OspA and B were the major components of the soluble fraction obtained after Triton partitioning of strain B31. In contrast, when Triton extraction was carried out after trypsin-treatment, the OspB band is not seen. Further purification of OspA-B31 on a SP

15 Sepharose column resulted in a single band by SDS-PAGE. The yield following removal of detergent was approximately 2 mg per liter of culture. This method of purification of OspA, as described herein for strain B31, can be used for other isolates of Borrelia as well.

20 For strains such as strain K48, which lack OspB, trypsin treatment can be omitted.

Lipidation site of OspA-B31

¹⁴C-palmitic acid labeled OspA from strain B31 was purified as described above and partially digested with endoproteinase Asp-N (data not shown). Following digestion, a new band of lower molecular weight was apparent by SDS-PAGE, found by direct amino-terminal sequencing to begin at Asp₂₅. This band had no trace of radioactivity by autoradiography (data not shown). OspA and B contain a signal sequence (L-X-Y-C) similar to the consensus described for lipoproteins of E. coli, and it has been predicted that the lipidation site of OspA and B should be the amino-terminal cysteine (Brandt, M.E. et

PCT/US94/12352

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al., <u>Infect. Immun 58</u>: 983-991 (1990)). The results presented herein support this prediction.

Comparison of OspA Antibody Binding Regions in Nine <u>B.</u> Strains of Borrelia burgdorferi

The availability of the amino acid sequenced for OspA from a number of different isolates, combined with peptide mapping and Western blot analysis, permitted the identification of the antigenic domains recognized by monoclonal antibodies (MAbs) and allowed inference of 10 the key amino acid residues responsible for specific antibody reactivity.

Strains of Borrelia burgdorferi

Nine strains of Borrelia, including seven European strains and two North American strains, were used in 15 this study of antibody binding domains of several proteins. Information concerning the strains is summarized in Table I, below.

Table I. Representative Borrelia Strains

Table I. Representative Bolletia Strains					
Strain	Location and Source	Reference for Strain			
K48	Czechoslovakia, Ixodes ricinus	none			
PGau	Germany, human ACA	Wilske, B. et al., J. Clin. <u>Microbiol. 32</u> :340-350 (1993)			
DK29	Denmark, human EM	Wilske, B. et al.			
PKo	Germany, human EM	Wilske, B. et al.			
PTrob	Germany, human skin	Wilske, B. et al.			
Ip3	Khabarovsk, Russia, I. persulcatus	Asbrink, E. et al., <u>Acta</u> <u>Derm. Venereol. 64</u> : 506-512 (1984)			
Ip90	Khabarovsk, Russia, I. persulcatus	Asbrink, E. et al.			
25015	Millbrook, NY, I. persulcatus	Barbour, A.G. et al., <u>Curr.</u> <u>Microbiol.</u> 8:123-126 (1983)			
B31	Shelter Island, NY, I. scapularis	Luft, B.J. et al., <u>Infect.</u> <u>Immun. 60</u> : 4309-4321 (1992); ATCC 35210			
PKa1	Germany, human CSF	Wilske, B. et al.			
ZS7	Freiburg, Germany, I. ricinus	Wallich, R. et al., <u>Nucl.</u> <u>Acids Res. 17</u> : 8864 (1989)			
N40	Westchester Co., NY	Fikrig, E. et al., <u>Science</u> 250:553-556 (1990)			
PHei	Germany, human CSF	Wilske, B. et al.			
ACAI	Sweden, human ACA	Luft, B. J. et al., <u>FEMS</u> <u>Microbiol. Lett. 93</u> :73-68 (1992)			
PBo	Germany, human CSF	Wilske, B. et al.			

ACA = patient with acrodermatitis chronica atrophicans; EM = patient with erythema migrans; CSF = cerebrospinal fluid of patient with Lyme disease

Strains K48, PGau and DK29 were supplied by R. Johnson, University of Minnesota; PKo and pTrob were provided by B. Wilske and V. Preac-Mursic of the

Pettenkhofer Institute, Munich, Germany; and Ip3 and Ip90 were supplied by L. Mayer of the Center for Disease Control, Atlanta, Georgia. The North American strains included strain 25015, provided by J. Anderson of the Connecticut Department of Agriculture; and strain B31 (ATCC 35210).

Monoclonal Antibodies

Seven monoclonal antibodies (MAbs) were utilized in this study. Five of the MAbs (12, 13, 15, 83 and 336) were produced from hybridomas cloned and subcloned as previously 10 described (Schubach, W.H., et al., Infect. Immun. 59(6):1911-1915 (1991)). MAb H5332 (Barbour, A.G. et al., Infect. Immun. 41:795-804 (1983)) was a gift from Drs. Alan Barbour, University of Texas, and MAb CIII.78 (Sears, J.E. 15 et al., <u>J. Immunol. 147(6):1995-2000(1991)</u>) was a gift from Richard A. Flavell, Yale University. MAbs 12 and 15 were raised against whole sonicated B3; MAb 336 was produced against whole PGau; and MAbs 13 and 83 were raised to a truncated form of OspA cloned from the K48 strain and 20 expressed in E. coli using the T7 RNA polymerase system (McGrath, B.C. et al., Vaccines, Cold Spring Harbor Laboratory Press, Plainview, New York, pp. 365-370 (1993)). All MAbs were typed as being Immunoglobulin G (IgG).

Methods of Protein Cleavage, Western Blotting, and Amino-Terminal Sequencing

Prediction of the various cleavage sites was achieved by knowledge of the primary amino acid sequence derived from the full nucleotide sequences of OspA, many of which are currently available (see Table II, below). Cleavage sites can also be predicted based on the peptide sequence of OspA, which can be determined by standard techniques after isolation and purification of OspA by the method described above. Cleavage of several OspA isolates was

conducted to determine the localization of monoclonal antibody binding of the proteins.

Hydroxylamine-HCl (HA), N-chlorosuccinimide (NCS), and cyanogen bromide cleavage of OspA followed the methods

5 described by Bornstein (Biochem. 9 (12):2408-2421 (1970)), Shechter et al., (Biochem. 15 (23):5071-5075 (1976)), and Gross (in Hirs, C.H.W. (ed): Methods in Enzymology, (N.Y. Acad. Press), 11:238-255 (1967)) respectively. Protease cleavage by endoproteinase, Asp-N (Boehringer Mannheim,

10 Indianapolis, Indiana), was performed as described by Cleveland D.W. et al., (J. Biol. Chem. 252:1102-1106 (1977)). Ten micrograms of OspA were used for each reaction. The ratio of enzyme to OspA was approximately 1 to 10 (w/w).

Proteins and peptides generated by cleavage were separated by SDS-polyacrylamide gel electrophoresis (SDS-PAGE) (Laemmli, U.K., Nature (London) 227:680-685 (1970)), and electroblotted onto immobilon Polyvinylidine Difluoride (PVDF) membranes (Ploskal, M.G. et al., Biotechniques

20 4:272-283 (1986)). They were detected by amido black staining or by immunostaining with murine MAbs, followed by alkaline phosphatase-conjugated goat antimouse IgG. Specific binding was detected using a 5-bromo-4-chloro-3-indolylphosphate (BCIP)/nitroblue tetrazolium (NBT)

25 developer system (KPL Inc., Gathersburg, Maryland).

In addition, amino-terminal amino acid sequence analysis was carried out on several cleavage products, as described by Luft et al. (<u>Infect. Immun. 57</u>:3637-3645 (1989)). Amido black stained bands were excised from PVDF blots and sequenced by Edman degradation using a Biosystems model 475A sequenator with model 120A PTH analyzer and model 900A control/data analyzer.

25

Cleavage Products of Outer Surface Protein A Isolates Purified OspA-B31, labeled with 14C-palmitic acid, was fragmented with hydroxylamine-HCl (HA) into two peptides, designated HA1 and HA2 (data not shown). The HA1 band 5 migrated at 27 KD and retained its radioactivity, indicating that the peptide included the lipidation site at the N-terminus of the molecule (data not shown). From the predicted cleavage point, HA1 should correspond to residues 1 to 251 of OspA-B31. HA2 had a MW of 21.6 KD by SDS-PAGE. 10 with amino-terminal sequence analysis showing it to begin at Gly72, i.e. residues 72 to 273 of OspA-B31. contrast, HA cleaved OspA-K48 into three peptides, designated HA1, HA2, and HA3 with apparent MWs of 22KD, 16 KD and 12 KD, respectively. Amino-terminal sequencing 15 showed HA1 to start at Gly72, and HA3 at Gly142. found to have a blocked amino-terminus, as was observed for the full-length OspA protein. HA1, 2 and 3 of OspA-K48 were predicted to be residues 72-274, 1 to 141 and 142 to 274, respectively.

N-Chlorosuccinimide (NCS) cleaves tryptophan (W), which is at residue 216 of OspA-B31 or residue 217 of OspA-K48 (data not shown). NCS cleaved OspA-B31 into 2 fragments, NCS1, with MW of 23 KD, residues 1-216 of the protein, and NCS2 with a MW of 6.2 KD, residues 217 to 273 (data not shown). Similarly, K48 OspA was divided into 2 pieces, NCS1 residues 1-217, and NCS2 residues 218 to 274 (data not shown).

Cleavage of OspA by cyanogen bromide (CNBr) occurs at the carboxy side of methionine, residue 39. The major fragment, CNBr1, has a MW of 25.7 KD, residues 39-274 by amino-terminal amino acid sequence analysis (data not shown). CNBr2 (about 4 KD) could not be visualized by amido black staining; instead, lightly stained bands of about 20 KD MW were seen. These bands reacted with anti-

OspA MAbs, and most likely were degradation products due to cleavage by formic acid.

Determination of Antibody Binding Domains for Anti-OspA Monoclonal Antibodies

The cleavage products of OspA-B31 and OspA-K48 were analyzed by Western blot to assess their ability to bind to the six different MAbs. Preliminary Western blot analysis of the cleavage products demonstrated that strains K48 and DK29 have similar patterns of reactivity, as do IP3, PGau The OspA of strain PTrob was immunologically distinct from the others, being recognized only by MAb 336. MAb 12 recognized only the two North American strains, B31 and 25015. When the isolates were separated into genogroups, it was remarkable that all the MAbs, except MAb 12, crossed over to react with multiple genogroups. 15

MAb12, specific for OspA-B31, bound to both HA1 and HA2 of OspA-B31. However, cleavage of OspA-B31 by NCS at residue Trp216 created fragments which did not react with MAb12, suggesting that the relevant domain is near or is 20 structurally dependent upon the integrity of this residue (data not shown). MAb 13 bound only to OspA-K48, and to peptides containing the amino-terminus of that molecule (e.g. HA2; NCS1). It did not bind to CNBr1 residues 39 to Thus the domain recognized by MAb13 is in the aminoterminal end of OspA-K48, near Met38.

MAb15 reacts with the OspA of both the B31 and K48 strains, and to peptides containing the N-terminus of OspA, such as HA1 of OspA-B31 and NCS1, but not to peptides HA2 of OspA-B31 and HA1 of OspA-K48 (data not shown). peptides include residue 72 to the C-terminus of the molecules. MAb15 bound to CNBr1 of OspA-K48, indicating the domain for this antibody to be residues 39 to 72, specifically near Gly72 (data not shown).

MAD83 binds to OspA-K48, and to peptides containing the C-terminal portion of the molecule, such as HA1. They do not bind to HA2 of OspA-K48, most likely because the C-terminus of HA2 of OspA-K48 ends at 141. Similar to MAD12 and OspA-B31, binding of MADS 83 and CIII.78 is eliminated by cleavage of OspA at the tryptophan residue. Thus binding of MADS 12, 83 and CIII.78 to OspA depends on the structural integrity of the Trp216 residue, which appears to be critical for antigenicity. Also apparent is that, although these MADS bind to a common antigenic domain, the precise epitopes which they recognize are distinct from one another given the varying degrees of cross-reactivity to these MADS among strains.

Although there is similar loss of binding activity of

MAb336 with cleavage at Trp216, this MAb does not bind to
HA1 of OspA-B31, suggesting the domain for this antibody
includes the carboxy-terminal end of the molecule,
inclusive of residues 251 to 273. Low MW peptides, such as
HA3 (10 KD) and NCS2 (6KD), of OspA-K48 do not bind this

MAb on Western blots. In order to confirm this
observation, we tested binding of the 6 MAbs with a
recombinant fusion construct p3A/EC that contains a trpE
leader protein fused with residues 217 to 273 of OspA-B31
(Schubach, W.H. et al., Infect. Immun. 59(6): 1911-1915

(1991)). Only MAb336 reacted with this construct (data not
shown). Peptides and antigenic domains localized by
fragmentation of OspA are summarized in Figure 1.

Mapping of Domains to Define the Molecular Basis for the Serotype Analysis

To define the molecular basis for the serotype analysis of OspA, we compared the derived amino acid sequences of OspA for the nine isolates (Figure 2). At the amino terminus of the protein, these predictions can be more precise given the relatively small number of amino

acid substitutions in this region compared to the carboxy Domain 1, which is recognized by MAb13, includes residues Leu34 to Leu41. MAb13 only binds to the OspA of species K48, DK29 and IP90. Within this region, residue 37 5 is variable, however Gly37 is conserved amongst the three reactive strains. When Gly37 is changed to Glu37, as it is in OspA of strains B31, pTrob, PGau, and PKo, MAb13 does not recognize the protein (data not shown). By similar analysis, it can be seen that Asp70 is a crucial residue 10 for Domain 2, which includes residues 65 to 75 and is recognized by MAb15. Domain 3 is reactive with MAbs H5332, 12 and 83, and includes residues 190-220. It is clear that significant heterogeneity exists between MAbs reactive with this domain, and that more than one conformational epitope 15 must be contained within the sequence. Domain 4 binds MAb336, and includes residues 250 to 270. In this region, residue 266 is variable and therefore may be an important determinant. It is apparent, however, that other determinants of the reactivity of this monoclonal antibody 20 reside in the region comprising amino acids 217-250. Furthermore, the structural integrity of Trp216 is essential for antibody reactivity in the intact protein. Finally, it is important to stress that Figure 2 indicates only the locations of the domains, and does not necessarily 25 encompass the entire domain. Exact epitopes are being analyzed by site-directed mutagenesis of specific residues.

Overall, evidence suggests that the N-terminal portion is not the immunodominant domain of OspA, possibly by virtue of its lipidation, and the putative function of the lipid moiety in anchoring the protein to the outer envelope. The C-terminal end is immunodominant and includes domains that account in part for structural heterogeneity (Wilske, B. et al., Med. Microbiol. Immunol. 181: 191-207 (1992)), and may provide epitopes for antibody

neutralization (Sears, J.E. et al., J. Immunol. 147(6): 1995-2000 (1991)), and relate to other activities, such as the induction of T-cell proliferation (Shanafel, M.M., et al., J. Immunol. 148: 218-224 (1992)). There are common 5 epitopes in the carboxy-end of the protein that are shared among genospecies which may have immunoprotective potential (Wilske, B., et al., Med. Microbiol. Immunol. 181: 191-207 (1992)).

Prediction of secondary structure on the basis of 10 hydropathy analysis and circular dichroism and fluorescence spectroscopy measurements (McGrath, B.C., et al., Vaccines, Cold Spring Harbor Laboratory Press, Plainview, New York; pp. 365-370 (1993)) suggest domains 3 and 4 to be in a region of the molecule with a propensity to form alpha-15 helix, whereas domains 1 and 2 occur in regions predicted to be beta-sheets (see Figure 1). These differences may distinguish domains in accessibility to antibody or to reactive T-cells (Shanafel, M.M. et al., J. Immunol. 148: 218-224 (1992)). Site-directed mutagenesis of specific 20 epitopes, as described below in Example 2, aids in identifying exact epitopes.

Identification of an Immunologically Example 2. Important Hypervariable Domain of the Major Outer Surface Protein A of Borrelia

This Example describes epitope mapping studies using chemically cleaved OspA and TrpE-OspA fusion proteins. studies indicate a hypervariable region surrounding the single conserved tryptophan residue of OspA (at residue 216, or in some cases 217), as determined by a moving 30 window population analysis of OspA from fifteen European and North American isolates of Borrelia. The hypervariable region is important for immune recognition.

Site-directed mutagenesis was also conducted to examine the hypervariable regions more closely. Fluorescence and circular dichroism spectroscopy have indicated that the conserved tryptophan is part of an 5 alpha-helical region in which the tryptophan is buried in a hydrophobic environment (McGrath, B.C., et al., Vaccines, Cold Spring Harbor Laboratory Press, Plainview, New York; pp. 365-370 (1993)). More polar amino acid side-chains flanking the tryptophan are likely to be exposed to the hydrophilic solvent. The hypervariability of these solvent-exposed residues among the various strains of Borrelia suggested that these amino acid residues may contribute to the antigenic variation in OspA. Therefore, site-directed mutagenesis was performed to replace some of 15 the potentially exposed amino acid side chains in the protein from one strain with the analogous residues of a second strain. The altered proteins were then analyzed by Western Blot using monoclonal antibodies which bind OspA on the surface of the intact, non-mutated spirochete. 20 results indicated that certain specific amino acid changes near the tryptophan can abolish reactivity of OspA to these monoclonal antibodies.

A. <u>Verification of Clustered Polymorphisms in Outer</u> <u>Surface Protein A Sequences</u>

Cloning and sequencing of the OspA protein from fifteen European and North American isolates (described above in Table I) demonstrated that amino acid polymorphism is not randomly distributed throughout the protein; rather, polymorphism tended to be clustered in three regions of OspA. The analysis was carried out by plotting the moving, weighted average polymorphism of a window (a fixed length subsection of the total sequence) as it is slid along the sequence. The window size in this analysis was thirteen amino acids, based upon the determination of the largest

number of significantly deviating points as established by the method of Tajima (<u>J. Mol. Evol. 33</u>: 470-473 (1991)). The average weighted polymorphism was calculated by summing the number of variant alleles for each site. Polymorphism calculations were weighted by the severity of amino acid replacement (Dayhoff, M.O. et al., in: Dayhoff, M.O. (ed.) Atlas of Protein Sequence and Structure NBRF, Washington, Vol. 5, Suppl. 3: 345 (1978)). The sum was normalized by the window size and plotted. The amino acid sequence 10 position corresponds to a window that encompasses amino acids 1 through 13. Bootstrap resampling was used to generate 95% confidence intervals on the sliding window analysis. Since Borrelia has been shown to be clonal, the bootstrap analysis should give a reliable estimate of the 15 expected variance out of polymorphism calculations. bootstrap was iterated five hundred times at each position, and the mean was calculated from the sum of all positions. The clonal nature of Borrelia ensures that the stochastic variance that results from differing genealogical histories of the sequence positions (as would be expected if recombination were prevalent) will be minimized.

This test verified that the three regions around the observed peaks all have significant excesses of polymorphism. Excesses of polymorphism were observed in 25 the regions including amino acid residues 132-145, residues 163-177, and residues 208-221 (Figure 3). An amino acid alignment between residues 200 and 220 for B31, K48 and the four site-directed mutants is shown in Figure 4. The amino acid 208-221 region includes the region of OspA which has 30 been modeled as an oriented alpha-helix in which the single tryptophan residue at amino acid 216 is buried in a hydrophobic pocket, thereby exposing more polar amino acids to the solvent (Figure 5) (France, L.L., et al., Biophys. Acta 1120: 59 (1992)). These potentially solventexposed residues showed considerable variability among the

OspAs from various strains and may be an important component of OspA antigenic variation. For the purposes of generating chimeric proteins, the hypervariable domains of interest are Domain A, which includes amino acid residues 120-140 of OspA; Domain B, which includes residues 150-180; and Domain C, which includes residues 200-216 or 217.

B. Site-Directed Mutagenesis of the Hypervariable Region
Site-directed mutagenesis was performed to convert
residues within the 204-219 domain of the recombinant B31

10 OspA to the analogous residues of a European OspA variant,
K48. In the region of OspA between residues 204 and 219,
which includes the helical domain (amino acids 204-217),
there are seven amino acid differences between OspA-B31 and
OspA-K48. Three oligonucleotides were generated, each
15 containing nucleotide changes which would incorporate K48
amino acids at their analogous positions in the B31 OspA
protein. The oligos used to create the site-directed
mutants were:

5'-CTTAATGACTCTGACACTAGTGC-3' (#613, which converts

20 threonine at position 204 to serine, and serine at 206 to
threonine (Thr204-Ser, Thr206-Ser)) (SEQ ID NO. 1);

5'-GCTACTAAAAAAACCGGGAAATGGAATTCA-3' (#625, which converts
alanine at 214 to glycine, and alanine at 215 to lysine
(Ala214-Gly, Ala215-Lys)) (SEQ ID NO. 2); and

5'-GCAGCTTGGGATTCAAAAACATCCACTTTAACA-3' (#640, which
converts asparagine at 217 to aspartate, and glycine at
219 to lysine (Asn217-Asp, Gly219-Lys)) (SEQ ID NO. 3).

Site-directed mutagenesis was carried out by performing mutagenesis with pairs of the above oligos.

Three site-directed mutants were created, each with two changes: OspA 613 (Thr204-Ser, Thr206-Ser), OspA 625 (Ala214-Gly, Ala215-Lys), and 640 (Asn217-Asp, Gly219-Lys). There were also two proteins with four changes: OspA 613/625 (Thr204-Ser, Thr206-Ser, Ala214-Gly, Ala215-Lys)

and OspA 613/640 (Thr204-Ser, Thr206-Ser, Asn217-Asp, Gly219-Lys).

Specificity of Antibody Binding to Epitopes of the Non-mutated Hypervariable Region

Monoclonal antibodies that agglutinate spirochetes, including several which are neutralizing in vitro, recognize epitopes that map to the hypervariable region around Trp216 (Barbour, A.G. et al., Infect. and Immun. 41: 759 (1983); Schubach, W.H. et al., Infect. and Immun. 59: 1911 (1991)). Western Blot analysis demonstrated that

chemical cleavage of OspA from the B31 strain at Trp 216 abolishes reactivity of the protein with the agglutinating Mab 105, a monoclonal raised against B31 spirochetes (data not shown). The reagent, n-chlorosuccinimide (NCS),

15 cleaves OspA at the Trp 216, forming a 23.2kd fragment and a 6.2kd peptide which is not retained on the Imobilon-P membrane after transfer. The uncleaved material binds Mab 105; however, the 23.2kd fragment is unreactive. Similar Western blots with a TrpE-OspA fusion protein containing

20 the carboxy-terminal portion of the OspA protein demonstrated that the small 6.2kd piece also fails to bind Mab 105 (Schubach, W.H. et al., <u>Infect. and Immun. 59</u>: 1911 (1991)).

Monoclonal antibodies H5332 and H3TS (Barbour, A.G. et al., Infect. and Immun. 41: 759 (1983)) have been shown by immunofluorescence to decorate the surface of fixed spirochetes (Wilske, B. et al., World J. Microbiol. 7: 130 (1991)). These monoclonals also inhibit the growth of the organism in culture. Epitope mapping with fusion proteins has confirmed that the epitopes which bind these Mabs are conformationally determined and reside in the carboxy half of the protein. Mab H5332 is cross-reactive among all of the known phylogenetic groups, whereas Mab H3TS and Mab 105 seem to be specific to the B31 strain to which they were

raised. Like Mab 105, the reactivities of H5332 and H3TS to OspA are abrogated by fragmentation of the protein at Trp216 (data not shown). Mab 336 was raised to whole spirochetes of the strain P/Gau. It cross-reacts to OspA from group 1 (the group to which B31 belongs) but not to group 2 (of which K48 is a member). Previous studies using fusion proteins and chemical cleavage have indicated that this antibody recognizes a domain of OspA in the region between residues 217 and 273 (data not shown). All of these Mabs will agglutinate the B31 spirochete.

Western Blot Analysis of Antibody Binding to Mutated Hypervariable Regions

Mabs were used for Western Blot analysis of the sitedirected OspA mutants induced in E.coli using the T7

15 expression system (Dunn, J.J. et al., Protein Expression
and Purification 1: 159 (1990)). E. coli cells carrying
Pet9c plasmids having a site-directed OspA mutant insert
were induced at mid-log phase growth with IPTG for four
hours at 37°C. Cell lysates were made by boiling an
20 aliquot of the induced cultures in SDS gell loading dye,
and this material was then loaded onto a 12% SDS gell
(BioRad mini-Protean II), and electrophoresed. The
proteins were then transferred to Imobilon-P membranes
(Millipore) 70V, 2 hour at 4°C using the BioRad mini
25 transfer system. Western analysis was carried out as
described by Schubach et al. (Infect. Immun. 59: 1911
(1991)).

Western Blot analysis indicated that only the 625 mutant (Ala214-Gly and Ala215-Lys) retained binding to the agglutinating monoclonal H3TS (data not shown). However, the 613/625 mutant which has additional alterations to the amino terminus of Trp216 (Ser204-Thr and Thr206-Ser) did not bind this monoclonal. Both 640 and 613/640 OspAs which have the Asn217-Asp and Gly219-Lys changes on the carboxy-

terminal side of Trp216 also failed to bind Mab H3TS. indicated that the epitope of the B31 OspA which binds H3TS is comprised of amino acid side-chains on both sides of Trp216.

The 613/625 mutant failed to bind Mabs 105 and H5332, while the other mutants retained their ability to bind these Mabs. This is important in light of the data using fusion proteins that indicate that Mab 105 behaves more like Mab H3TS in terms of its serotype specificity and 10 binding to OspA (Wilske, B. et al., Med. Microbiol. Immunol. 181: 191 (1992)). The 613/625 protein has, in addition to the differences at residues Thr204 and Ser206, changes immediately amino-terminal to Trp216 (Ala214-Gly and Ala215-Lys). The abrogation of reactivity of Mabs 105 and H5332 to this protein indicated that the epitopes of OspA which bind these monoclonals are comprised of residues on the amino-terminal side of Trp216.

The two proteins carrying the Asn217-Asp and Gly219-Lys replacements on the carboxy-terminal side of Trp216 20 (OspAs 640 and 613/640) retained binding to Mabs 105 and H5332; however, they failed to react with Mab 336, a monoclonal which has been mapped with TrpE-OspA fusion proteins and by chemical cleavage to a more carboxyterminal domain. This result may explain why Mab 336 25 failed to recognize the K48-type of OspA (Group 2).

It is clear that amino acids Ser204 and Thr206 play an important part in the agglutinating epitopes in the region of the B31 OspA flanking Trp216. Replacement of these two residues altered the epitopes of OspA that bind Mabs 105, 30 H3TS and H5332. The ability of the 640 changes alone to abolish reactivity of Mab 336 indicated that Thr204 and Ser206 are not involved in direct interaction with Mab 336.

The results indicated that the epitopes of OspA which are available to Mabs that agglutinate spirochetes are 35 comprised at least in part by amino acids in the immediate vicinity of Trp216. Since recent circular dichroism analysis indicated that the structures of B31 and K48 OspA differ very little within this domain, it is unlikely that the changes made by mutation have radically altered the overall structure of the OspA protein (France, L.L. et al., Biochem. Biophys. Acta 1120: 59 (1992); and France et al., Biochem. Biophys Acta, submitted (1993)). This hypothesis is supported by the finding that the recombinant, mutant OspAs exhibit the same high solubility and purification properties as the parent B31 protein (data not shown).

In summary, amino acid side-chains at Ser204 and Thr206 are important for many of the agglutinating epitopes. However, a limited set of conservative changes at these sites were not sufficient to abolish binding of all of the agglutinating Mabs. These results suggested that the agglutinating epitopes of OspA are distinct, yet may have some overlap. The results also supported the hypothesis that the surface-exposed epitope around Trp216 which is thought to be important for immune recognition and neutralization is a conformationally-determined and complex domain of OspA.

EXAMPLE 3. Borrelia Strains and Proteins

Proteins and genes from any strain of Borrelia can be utilized in the current invention. Representative strains are summarized in Table I, above.

A. Genes Encoding Borrelia Proteins

The chimeric peptides of the current invention can comprise peptides derived from any Borrelia proteins.

Representative proteins include OspA, OspB, OspC, OspD,

pl2, p39, p41 (fla), p66, and p93. Nucleic acid sequences encoding several Borrelia proteins are presently available (see Table II, below); alternatively, nucleic acid

sequences encoding *Borrelia* proteins can be isolated and characterized using methods such as those described below.

Table II. References for Nucleic Acid Sequences for Several Proteins of Various Borrelia Strains

Strai n	р93	OspA	p41 (fla)
K48	X69602 (SID 67)	X62624 (SID 8)	X69610 (SID 49)
PGau	SID 73	X62387 (SID 10)	X69612 (SID 51)
DK29		X63412 (SID 137)	X69608 (SID 53)
PKo	X69803 (SID 77)	X65599 (SID 141)	X69613 (SID 131)
PTrob	X69604 (SID 71)	X65598 (SID 135)	X69614 (SID 55)
Ip3	•	X70365 (SID 140)	-
Ip90	ND	Kryuchechnikov, V.N. et al., <u>J.Microbiol.</u> <u>Epid. Immunobiol.</u> 12:41-44 (1988) (SID 138)	-
25015	X70365 (SID 75)	Fikrig, E.S. et al., <u>J. Immunol. 7</u> :2256- 2260 1992) SID 12)	-
B31	Perng, G.C. et al., <u>Infect.</u> <u>Immun. 59:</u> 2070- 74 (1992); Luft, B.J. et al., <u>Infect.</u> <u>Immun. 60:</u> 4309- 4321 (1992) (SID 65)	Bergstrom, S. et al., Mol. Microbiol. 3:479-486 (1989) (SID 6)	Gassmann, G.S. et al., <u>Nucl.</u> <u>Acids Res. 17</u> : 3590 (1989) (SID 127)
PKal	-	X69606 (SID.132)	X69611 (SID 129)
ZS7		Jonsson, M. et al., <u>Infect. Immun.</u> <u>60</u> :1845-1853 (1992) (SID 134)	-
N40		Kryuchechnikov, V.N. et al. (SID 133)	-
PHei	-	X65600 (SID 136)	-
ACAI	-	Kryuchechnikov, V.N. et al. (SID 142)	-
PBo	X69601 (SID 69)	X65605 (SID 139)	X69610 (SID 130)

Numbers with an "X" prefix are GenBank data base accession numbers. SID = SEQ ID NO.

B. Isolation of Borrelia Genes

Nucleic acid sequences encoding full length, lipidated proteins from known Borrelia strains were isolated using the polymerase chain reaction (PCR) as described below. In 5 addition, nucleic acid sequences were generated which encoded truncated proteins (proteins in which the lipidation signal has been removed, such as by eliminating the nucleic acid sequence encoding the first 18 amino acids, resulting in non-lipidated proteins). 10 proteins were generated which encoded polypeptides of a particular gene (i.e., encoding a segment of the protein which has a different number of amino acids than the protein does in nature). Using similar methods as those described below, primers can be generated from known 15 nucleic acid sequences encoding Borrelia proteins and used to isolate other genes encoding Borrelia proteins. Primers can be designed to amplify all of a gene, as well as to amplify a nucleic acid sequence encoding truncated protein sequences, such as described below for OspC, or nucleic 20 acid sequences encoding a polypeptide derived from a Borrelia protein. Primers can also be designed to incorporate unique restriction enzyme cleavage sites into the amplified nucleic acid sequences. Sequence analysis of the amplified nucleic acid sequences can then be performed 25 using standard techniques.

Cloning and Sequencing of OspA Genes and Relevant Nucleic Acid Sequences

Borrelia OspA sequences were isolated in the following manner: 100 μ l reaction mixtures containing 50 mM KCl, 10 mM TRIS-HCl (pH 8,3), 1.5 mM MgCl₂, 200 μ M each NTP, 2.5 units of TaqI DNA polymerase (Amplitaq, Perkin-Elmer/Cetus) and 100 pmol each of the 5' and 3' primers (described below) were used. Amplification was performed in a Perkin-Elmer/Cetus thermal cycler as described (Schubach, W.H. et

al., Infect. Immun. 59:1811-1915 (1991)). The amplicon was visualized on an agarose gel by ethidium bromide staining. Twenty nanograms of the chloroform-extracted PCR product were cloned directly into the PC-TA vector (Invitrogen) by following the manufacturer's instructions. Recombinant colonies containing the amplified fragment were selected, the plasmids were prepared, and the nucleic acid sequence of each OspA was determined by the dideoxy chaintermination technique using the Sequenase kit (United States Biochemical). Directed sequencing was performed with M13 primers followed by OspA-specific primers derived from sequences, previously obtained with M13 primers.

Because the 5' and 3' ends of the OspA gene are highly conserved (Fikrig, E.S. et al., J. Immunol. 7:2256-2260

15 (1992); Bergstrom, S. et al., Mol. Microbiol. 3: 479-486 (1989); Zumstein, G. et al., Med. Microbiol. Immunol. 181: 57-70 (1992)), the 5' and 3' primers for cloning can be based upon any known OspA sequences. For example, the following primers based upon the OspA nucleic acid sequence 20 from strain B31 were used:

5'-GGAGAATATTATGAAA-3' (-12 to +6) (SEQ ID NO. 4); and 5'-CTCCTTATTTTAAAGCG-3' (+826 to +809) (SEQ ID NO. 5). (Schubach, W.H. et al., <u>Infect. Immun 59</u>:1811-1915 (1991)).

OspA genes isolated in this manner include those for strains B31, K48, PGau, and 25015; the nucleic acid sequences are depicted in the sequence listing as SEQ ID NO. 6 (OspA-B31), SEQ ID NO. 8 (OspA-K48), SEQ ID NO. 10 (OspA-PGau), and SEQ ID NO. 12 (OspA-25015). An alignment of these and other OspA nucleic acid sequences is shown in Figure 42. The amino acid sequences of the proteins encoded by these nucleic acid sequences are represented as SEQ ID NO. 7 (OspA-B31), SEQ ID NO. 9 (OspA-K48), SEQ ID NO. 11 (OspA-PGau), and SEQ ID NO. 13 (OspA-25015).

The following primers were used to generate specific nucleic acid sequences of the OspA gene, to be used to

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generate chimeric nucleic acid sequences (as described in Example 4):

- 5'-GTCTGCAAAAACCATGACAAG-3' (plus strand primer #369) (SEQ ID NO. 14);
- 5 5'-GTCATCAACAGAAGAAAATTC-3' (plus strand primer #357) (SEQ ID NO 15);
 - 5'-CCGGATCCATATGAAAAATATTTATTGGG-3' (plus strand primer #607) (SEQ ID NO. 16);
 - 5'-CCGGGATCCATATGGCTAAGCAAAATGTTAGC-3' (plus strand primer
- 10 #584) (SEQ ID NO. 17); 5'-GCGTTCAAGTACTCCAGA-3' (minus strand primer #200) (SEQ ID NO. 18);
 - 5'-GATATCTAGATCTTATTTTAAAGCGTT-3' (minus strand primer #586) (SEQ ID NO. 19); and
- 15 5'-GGATCCGGTGACCTTTTAAAGCGTTTTTAAT-3' (minus strand primer #1169) (SEQ ID NO. 20).

Cloning and Sequencing of OspB

Similar methods were also used to isolate OspB genes. One OspB genes isolated is represented as SEQ ID NO. 21 (OspB-B31); its encoded amino acid sequence is SEQ ID NO. 22.

The following primers were used to generate specific nucleic acid sequences of the OspB gene, to be used in generation of chimeric nucleic acid sequences (see Example

25 4):

20

- 5'-GGTACAATTACAGTACAA-3' (plus strand primer #721) (SEQ ID NO. 23);
- 5'-CCGAGAATCTCATATGGCACAAAAAGGTGCTGAGTCAATTGG-3' (plus strand primer #1105) (SEQ ID NO. 24);
- 30 5'-CCGATATCGGATCCTATTTTAAAGCGTTTTTAAGC-3' (minus strand
 primer # 1106) (SEQ ID NO. 25); and
 5'-GGATCCGGTGACCTTTTAAAGCGTTTTTAAG-3' (minus strand primer
 #1170) (SEQ ID NO. 26).

Cloning and Sequencing of OspC

Similar methods were also used to isolate OspC genes. The following primers were used to isolate entire OspC genes from Borrelia strains B31, K48, PKO, and pTrob:

5'-GTGCGCGACCATATGAAAAAGAATACATTAAGTGCG-3' (plus strand primer having Ndel site combined with start codon) (SEQ ID NO. 27), and

5'-GTCGGCGGATCCTTAAGGTTTTTTTGGACTTTCTGC-3' (minus strand primer having BamHl site followed by stop codon) (SEQ ID NO. 28).

The nucleic acid sequences of the OspC genes were then determined by the dideoxy chain-termination technique using the Sequenase kit (United States Biochemical). OspC genes isolated and sequenced in this manner include those for strains B31, K48, PKo, and Tro; the nucleic acid sequences are depicted in the sequence listing as SEQ ID NO. 29 (OspC-B31), SEQ ID NO. 31 (OspC-K48), SEQ ID NO. 33 (OspC-PKo), and SEQ ID NO. 35 (OspC-Tro). An alignment of these sequences is shown in Figure 38. The amino acid sequences of the proteins encoded by these nucleic acid sequences are represented as SEQ ID NO. 30 (OspC-B31), SEQ ID NO. 32 (OspC-K48), SEQ ID NO. 34 (OspC-PKo), and SEQ ID NO. 36 (OspC-Tro).

Truncated OspC genes were generated using other

25 primers. These primers were designed to amplify nucleic acid sequences, derived from the OspC gene, that lacked the nucleic acids encoding the signal peptidase sequence of the full-length protein. The primers corresponded to bp 58-75 of the natural protein, with a codon for Met-Ala attached

30 ahead. For strain B31, the following primer was used:

5'-GTGCGCGACCATATGGCTAATAATTCAGGGAAAGAT-3' (SEQ ID NO.

37).

For strain PKo,

5'-GTGCGCGACCATATGGCTAGTAATTCAGGGAAAGGT-3' (SEQ ID NO. 38)
35 was used.

For strains pTrob and K48, 5'-GTGCGCGACCATATGGCTAATAATTCAGGTGGGGAT-3' (SEQ ID NO. 39) was used.

Additional primers were also designed to amplify

nucleic acids encoding particular polypeptides, for use in
creation of chimeric nucleic acid sequences (see Example

4). These primers included:

5'-CTTGGAAAATTATTTGAA-3' (plus strand primer #520) (SEQ ID NO. 40);

5'-CACGGTCACCCCATGGGAAATAATTCAGGGAAAGG-3' (plus strand primer #58) (SEQ ID NO. 41);

5'-TATAGATGACAGCAACGC-3' (minus strand primer #207) (SEQ ID NO. 42); and

5'-CCGGTGACCCCATGGTACCAGGTTTTTTTGGACTTTCTGC-3' (minus strand primer #636) (SEQ ID NO. 43).

Cloning and Sequencing of OspD

Similar methods can be used to isolate OspD genes. An alignment of four OspD nucleic acid sequences (from strains pBo, PGau, DK29, and K48) is shown in Figure 39.

20 Cloning and Sequencing of p12

The p12 gene was similarly identified. Primers used to clone the entire p12 gene included: 5'CCGGATCCATATGGTTAAAAAAATAATATTTTTTC-3' (forward primer # 757) (SEQ ID NO. 44); and 5'-

25 GATATCTAGATCTTTAATTGCTCTGCTCACTCTCTC-3' (reverse primer #758) (SEQ ID NO. 45).

To amplify a truncated p12 gene (one in which the transcribed protein is non-lipidated, and begins at amino acid 18 of the native sequence), the following primers were used: 5'-CCGGGATCCATATGGCTAGTGCAATTGGTCGTGG-3' (forward primer # 759) (SEQ ID NO. 46); and primer #758 (SEQ ID NO. 45).

Cloning and Sequencing of p41 (fla)

A similar approach was used to clone and sequence genes encoding the p41 (fla) protein. The p41 sequences listed in Table II with GenBank accession numbers were 5 isolated using the following primers from strain B31: 5'-ATGATTATCAATCATAAT-3' (+1 to +18) (SEQ ID NO. 47); and 5'-TCTGAACAATGACAAAAC-3' (+1008 to +991) (SEQ ID NO. 48). The nucleic acid sequences of p41 isolated in this manner are depicted in the sequence listing as SEQ ID NO. 51 (p41-10 PGau), and SEQ ID NO. 53 (p41-DK29). An alignment of several p41 nucleic acid sequences, including those for strains B31, pKa1, PGau, pBo, DK29, and pKo, is shown in Figure 41. The amino acid sequences of the proteins encoded by these nucleic acid sequences are represented as 15 SEQ ID NO. 50 (p41-K48), SEQ ID NO. 52 (p41-PGau), SEQ ID NO. 54 (p41-DK29), SEQ ID NO. 56 (p41-PTrob), and SEQ ID NO. 58 (p41-PHei).

Other primers were designed to amplify nucleic acid sequences encoding polypeptides of p41, to be used in chimeric nucleic acid sequences. These primers included: 5'-TTGGATCCGGTCACCCCATGGCTCAATATAACCAATG-3' (minus strand primer #122) (SEQ ID NO. 59);

- 5'-TTGGATCCGGTCACCCCATGGCTTCTCAAAATGTAAG-3' (plus strand primer # 140) (SEQ ID NO. 60);
- 25 5'-TTGGATCCGGTGACCAACTCCGCCTTGAGAAGG-3' (minus strand primer # 234) (SEQ ID NO. 61); and
 - 5'-TTGGATCCGGTGACCTATTTGAGCATAAGATGC-3' (minus strand primer #141) (SEQ ID NO. 62).

Cloning and Sequencing of p93

The same approach was also used to clone and sequence p93 protein. Genes encoding p93, as listed in Table II with GenBank accession numbers, were isolated by this method with the following primers from strain B31:

- 5'-GGTGAATTTAGTTGGTAAGG-3' (-54 to -35) (SEQ ID NO. 63); and
- 5'-CACCAGTTTCTTTAAGCTGCTCCTGC-3' (+1117 to +1092) (SEQ ID NO. 64).
- The nucleic acid sequences of p93 isolated in this manner are depicted in the sequence listing as SEQ ID NO. 65 (p93-B31), SEQ ID NO. 67 (p93-K48) SEQ ID NO. 69 (p93-PBo), SEQ ID NO. 71 (p93-PTrob), SEQ ID NO. 73 (p93-PGau), SEQ ID NO. 75 (p93-25015), and SEQ ID NO. 77 (p93-PKo).
- The amino acid sequences of the proteins encoded by these nucleic acid sequences are represented as SEQ ID NO. 66 (p93-B31), SEQ ID NO. 68 (p93-K48) SEQ ID NO. 70 (p93-PBO), SEQ ID NO. 72 (p93-PTrob), SEQ ID NO. 74 (p93-PGau), SEQ ID NO. 76 (p93-25015), and SEQ ID NO. 78 (p93-PKO).
- Other primers were used to amplify nucleic acid sequences encoding polypeptides of p93 to be used in generating chimeric nucleic acid sequences. These primers included:
- 5'-CCGGTCACCCCATGGCTGCTTTAAAGTCTTTA-3' (plus strand primer 20 #475) (SEQ ID NO. 79);
 - 5'-CCGGTCACCCCATGAATCTTGATAAAGCTCAG-3' (plus strand primer #900) (SEO ID NO. 80);
 - 5'-CCGGTCACCCCATGGATGAAAAGCTTTTAAAAAGT-3' (plus strand primer #1168) (SEQ ID NO. 81);
- 25 5'-CCGGTCACCCCCATGGTTGAGAAATTAGATAAG-3' (plus strand primer #1423) (SEQ ID NO. 82); and 5'-TTGGATCCGGTGACCCTTAACTTTTTTAAAG-3' (minus strand
 - primer # 2100) (SEQ ID NO. 83).

C. Expression of Proteins from Borrelia Genes

The nucleic acid sequences described above can be incorporated into expression plasmids, using standard techniques, and transfected into compatible host cells in order to express the proteins encoded by the nucleic acid

sequences. As an example, the expression the pl2 gene and the isolation of pl2 protein is set forth.

Amplification of the p12 nucleic acid sequence was conducted with primers that included a NdeI restriction 5 site into the nucleic acid sequence. The PCR product was extracted with phenol/chloroform and precipitated with ethanol. The precipitated product was digested and ligated into an expression plasmid as follows: 15 μ l (approximately 1 μ g) of PCR DNA was combined with 2 μ l 10X 10 restriction buffer for NdeI (Gibco/BRL), 1 μ l NdeI (Gibco/BRL), and 2 μ l distilled water, and incubated overnight at 37°C. This mixture was subsequently combined with 3 μ l 10X buffer (buffer 3, New England BioLabs), 1 μ l BamHI (NEB), and 6 μ l distilled water, and incubated at 37° 15 for two hours. The resultant material was purified by preparative gel electrophoresis using low melting point agarose, and the band was visualized under long wave ultraviolet light and excised from the gel. The gel slice was treated with Gelase using conditions recommended by the 20 manufacturer (Epicentre Technologies). The resulting DNA pelled was resuspended in 25-50 μl of 10 mM TRIS-CL (pH 8.0) and 1 mM EDTA (TE). An aliquot of this material was ligated into the Pet9c expression vector (Dunn, J. J. et al., Protein Expression and Purification 1: 159 (1990)).

To ligate the material into the Pet9c expression vector, 20-50 ng of pl2 nucleic acid sequences cut and purified as described above was combined with 5 μ l 10 One-Phor-All (OPA) buffer (Pharmacia), 30-60 ng Pet9c cut with NdeI and BamHI, 2.5 μ l 20 mM ATP, 2 μ l T4 DNA ligase 30 (Pharmacia) diluted 1:5 in 1X OPA buffer, and sufficient distilled water to bring the final volume to 50 μ l. mixture was incubated at 12°C overnight.

The resultant ligations were transformed into competent DH5-alpha cells and plated on nutrient agar 35 plates containing 50 μ g/ml kanamycin and incubated

overnight at 37 °C. DH5-alpha is used as a "storage strain" for T7 expression clones, because it is RecA deficient, so that recombination and concatenation are not problematic, and because it lacks the T7 RNA polymerase gene necessary to express the cloned gene. The use of this strain allows for cloning of potentially toxic gene products while minimizing the chance of deletion and/or rearrangement of the desired genes. Other cell lines having similar properties may also be used.

Kanamycin resistant colonies were single-colony purified on nutrient agar plates supplemented with kanamycin at 50 μ g/ml. A colony from each isolate was inoculated into 3-5 ml of liquid medium containing 50 μ g/ml kanamycin, and incubated at 37°C without agitation.

Plasmid DNA was obtained from 1 ml of each isolate using a hot alkaline lysis procedure (Mantiatis, T. et al., Molecular Cloning: A Laboratory Manual, cold Spring Harbor Laboratory, Cold Spring Harbor, NY (1982)).

Plasmid DNA was digested with EcoRI and BglII in the
following manner: 15 μl plasmid DNA was combined with 2 μl
10X buffer 3 (NEB), 1 μ EcoRI (NEB), 1 μl BglII (NEB) and 1
μl distilled water, and incubated for two hours at 37°C.
The entire reaction mixture was electrophoresed on an
analytical agarose gel. Plasmids carrying the p12 insert
were identified by the presence of a band corresponding to
925 base-pairs (full length p12) or 875 base-pairs
(nonlipidated p12).

One or two plasmid DNAs from the full length and nonlipidated p12 clones in Pet9c were used to transform

30 BL21 DE3 pLysS to kanamycin resistance as described by Studier et al. (Methods in Enzymology, Goeddel, D. (Ed.), Academic Press, 185: 60-89 (1990)). One or two transformants of the full length and nonlipidated clones were single-colony purified on nutrient plates containing

35 25 µg/ml chloramphenicol (to maintain pLysS) and 50 µg/ml

kanamycin at 37 °C. One colony of each isolate was inoculated into liquid medium supplemented with chloramphenicol and kanamycin and incubated overnight at The overnight culture was subcultured the following 5 morning into 500 ml of liquid broth with chloramphenicol (25 μ g/ml) and kanamycin (50 μ g/ml) and grown with aeration at 37°C in an orbital air-shaker until the absorbance at 600 nm reached 0.4-0.7. Isopropyl-thio-galactoside (IPTG) was added to a final concentration of 0.5 mM, for 10 induction, and the culture was incubated for 3-4 hours at 37° as before. The induced cells were pelleted by centrifugation and resuspended in 25 ml of 20 mM NaPO, (pH 7.7). A small aliquot was removed for analysis by gel electrophoresis. Expressing clones produced proteins which 15 migrated at the 12 kDa position.

A crude cell lysate was prepared from the culture as described for recombinant OspA by Dunn, J.J. et al., (Protein Expression and Purification 1: 159 (1990)). crude lysate was first passed over a Q-sepharose column (Pharmacia) which had been pre-equilibrated in Buffer A: 10 mM NaPO₄ (pH 7.7), 10 mM NaCl, 0.5 mM PMSF. The column was washed with 10 mM NaPO4, 50 mM NaCl and 0.5 mM PMSF and then pl2 was eluted in 10 mM NaPO4, 0.5 mM PMSF with a NaCl gradient from 50-400 mM. pl2 eluted approximately halfway 25 through the gradient between 100 and 200 mM NaCl. The peak fractions were pooled and dialyzed against 10 mM NaPo4 (pH 7.7), 10 mM NaCl, 0.5 mM PMSF. The protein was then concentrated and applied to a Sephadex G50 gel filtration column of approximately 50 ml bed volume (Pharmacia), in 10 30 mM NaPO₄, 200 mM NaCl, 0.5 mM PMSF. p12 would typically elute shortly after the excluded volume marker. fractions were determined by running small aliquots of all fractions on a gel. The p12 peak was pooled and stored in small aliquots at -20°C.

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Example 4. Generation of Chimeric Nucleic Acid Sequences and Chimeric Proteins

A. General Protocol for Creation of Chimeric Nucleic Acid Sequences

The megaprimer method of site directed mutagenesis and its modification were used to generate chimeric nucleic acid sequences (Sarkar and Sommer, Biotechniques 8(4): 404-407 (1990); Aiyar, A. and J. Leis, Biotechniques 14(3): 366-369 (1993)). A 5' primer for the first genomic template and a 3' fusion oligo are used to amplify the desired region. the fusion primer consists of a 3' end of the first template (DNA that encodes the amino-proximal polypeptide of the fusion protein), coupled to a 5' end of the second template (DNA that encodes the carboxy-proximal polypeptide of the fusion protein).

The PCR amplifications are performed using Taq DNA polymerase, 10X PCR buffer, and $MgCl_2$ (Promega Corp., Madison, WI), and Ultrapure dNTPs (Pharmacia, Piscataway, NJ). One μg of genomic template 1, 5 μ of 10 μM 5' oligo and 5 μl of 10 μM fusion oligo are combined with the following reagents at indicated final concentrations: 10X Buffer-Mg FREE (1X), $MgCl_2$ (2 mM), dNTP mix (200 μM each dNTP), Taq DNA polymerase (2.5 units), water to bring final volume to 100 μl . A Thermal Cycler (Perkin Elmer Cetus,

25 Norwalk, CT) is used to amplify under the following conditions: 35 cycles at 95°C for one minute, 55°C for two minutes, and 72° for three minutes. This procedure results in a "megaprimer".

The resulting megaprimer is run on a 1X TAE, 4% low-solution and the gelest and purified using the Promega Magic PCR Preps DNA purification system. Purified megaprimer is then used in a second PCR step. One μg of genomic template 2, approximately 0.5 μg of the megaprimer, and 5 μ of 10 μM 3'

oligo are added to a cocktail of 10X buffer, MgCl2, dNTPs and Tag at the same final concentrations as noted above, and brought to 100 μ l with water. PCR conditions are the same as above. The fusion product resulting from this 5 amplification is also purified using the Promega Magic PCR Preps DNA purification system.

The fusion product is then ligated into TA vector and transformed into E. coli using the Invitrogen (San Diego, CA) TA Cloning Kit. Approximately 50 ng of PCR fusion 10 product is ligated to 50 ng of pCRII vector with 1X Ligation Buffer, 4 units of T4 ligase, and brought to 10 Nl with water. This ligated product mixture is incubated at 12°C overnight (approximately 14 hours). Two $\mu \dot{1}$ of the ligation product mixture is added to 50 μl competent INC F' 15 cells and 2 μ beta mercaptoethanol. The cells are then incubated for 30 minutes, followed by heat shock treatment at 42°C for 60 seconds, and an ice quenching for two minutes. 450 μ l of warmed SOC media is then added to the cells, resulting in a transformed cell culture which is 20 incubated at 37°C for one hour with slight shaking. of the transformed cell culture is plated on LB + 50 $\mu g/\mu l$ ampicillin plates and incubated overnight at 37°C. white colonies are picked and added to individual overnight cultures containing 3 ml LB with ampicillin (50 μ g/ μ l).

The individual overnight cultures are prepared using Promega's Magic Miniprep DNA purification system. A small amount of the resulting DNA is cut using a restriction digest as a check. DNA sequencing is then performed to check the sequence of the fusion nucleic acid sequence, 30 using the United States Biochemical (Cleveland, OH) Sequenase Version 2.0 DNA sequencing kit. Three to five μg of plasmid DNA is used per reaction. 2 μ l 2M NaOH/2mM EDTA are added to the DNA, and the volume is brought to 20 μl The mixture is then incubated at room with water. temperature for five minutes. 7 μ l water, 3 μ l 3M NaAc, 75

 μ l EtOH are added. The resultant mixture is mixed by vortex and incubated for ten minutes at -70°C, and then subjected to microfugation. After microfuge for ten minutes, the supernatant is aspirated off, and the pellet 5 is dried in the speed vac for 30 second. 6 μ l water, 2 μ l annealing buffer, and 2 μl of 10 μM of the appropriate oligo is then added. This mixture is incubated for 10 minutes at 37°C and then allowed to stand at room temperature for 10 minutes. Subsequently, 5.5 μ l of label 10 cocktail (described above) is added to each sample of the mixture, which are incubated at room temperature for an additional five minutes. 3.5 μl labeled DNA is then added to each sample which is then incubated for five minutes at 37°C. 4 μ l stop solution is added to each well. 15 is denatured at 95° for two minutes, and then placed on ice.

Clones with the desired fusion nucleic acid sequences are then recloned in frame in the pEt expression system in the lipidated (full length) and non-lipidated (truncated, 20 i.e., without first 17 amino acids) forms. The product is amplified using restriction sites contained in the PCR primers. The vector and product are cut with the same enzymes and ligated together with T4 ligase. The resultant plasmid is transformed into competent E. coli using 25 standard transformation techniques. Colonies are screened as described earlier and positive clones are transformed into expression cells, such as E. coli BL21, for protein expression with IPTG for induction. The expressed protein in its bacterial culture lysate form and/or purified form 30 is then injected in mice for antibody production. The mice are bled, and the sera collected for agglutination, in vitro growth inhibition, and complement- dependent and independent lysis tests.

B. Specific Chimeric Nucleic Acid Sequences

Various chimeric nucleic acid sequences were The nucleic acid sequences are described as encoding polypeptides from Borrelia proteins. The chimeric 5 nucleic acid sequences are produced such that the nucleic acid sequence encoding one polypeptide is in the same reading frame as the nucleic acid sequence encoding the next polypeptide in the chimeric protein sequence encoded by the chimeric nucleic acid sequence. The proteins are 10 listed sequentially (in order of presence of the encoding sequence) in the description of the chimeric nucleic acid sequence. For example, if a chimeric nucleic acid sequence consists of bp 1-650 from OspA-1 and bp 651-820 from OspA-2 were sequenced, the sequence of the chimer would include 15 the first 650 base pairs from OspA-1 followed immediately by base pairs 651-820 of OspA-2.

OspA-K48/OspA-PGau A chimer of OspA from strain
K48 (OspA-K48) and OspA from strain PGau (OspA-PGau) was
generated using the method described above. This chimeric
nucleic acid sequence included bp 1-654 from OspA-K48,
followed by bp 655-820 from OspA-PGau. Primers used
included: the amino-terminal sequence of OspA primer #607
(SEQ ID NO. 16); the fusion primer,

5'-AAAGTAGAAGTTTTTGAATCCCATTTTCCAGTTTTTT-3' (minus strand primer #668-654) (SEQ ID NO. 84); the carboxy-terminal sequence of OspA primer #586 (SEQ ID NO. 19); and the sequence primers #369 (SEQ ID NO. 14) and #357 (SEQ ID NO. 15). The chimeric nucleic acid sequence is presented as SEQ ID NO. 85; the chimeric protein encoded by this chimeric nucleic acid sequence is presented as SEQ ID NO. 86.

OspA-B31/OspA-PGau A chimer of OspA from strain B31 (OspA-B31) and OspA from strain PGau (OspA-PGau) was generated

using the method described above. This chimeric nucleic acid sequence included bp 1-651 from OspA-B31, followed by bp 652-820 from OspA-PGau. Primers used included: the fusion primer,

5 5'-AAAGTAGAAGTTTTTGAATTCCAAGCTGCAGTTTT-3' (minus strand primer #668-651) (SEQ ID NO. 87); and the sequence primer, #369 (SEQ ID NO. 14). The chimeric nucleic acid sequence is presented as SEQ ID NO. 88; the chimeric protein encoded by this chimeric nucleic acid sequence is presented as SEQ ID NO. 89.

OspA-B31/OspA-K48 A chimer of OspA from strain B31 (OspA-B31) and OspA from strain K48 (OspA-K48) was generated using the method described above. This chimeric nucleic acid sequence included bp 1-651 from OspA-B31, followed by bp 652-820 from OspA-K48. Primers used included: the fusion primer,

5'-AAAGTGGAAGTTTTTGAATTCCAAGCTGCAGTTTTTTT-3' (minus strand primer #671-651) (SEQ ID NO. 90); and the sequence primer, #369 (SEQ ID NO. 14). The chimeric nucleic acid sequence is presented as SEQ ID NO. 91; the chimeric protein encoded by this chimeric nucleic acid sequence is presented as SEQ ID NO. 92.

OspA-B31/OspA-25015 A chimer of OspA from strain B31 (OspA-B31) and OspA from strain 25015 (OspA-25015) was generated using the method described above. This chimeric nucleic acid sequence included bp 1-651 from OspA-B31, followed by bp 652-820 from OspA-25015. Primers used included: the fusion primer, 5'-TAAAGTTGAAGTGCCTGCATTCCAAGCTGCAGTTT-3' (SEQ ID NO. 93). The chimeric nucleic acid sequence is presented as SEQ ID NO. 94; the chimeric protein encoded by this chimeric nucleic acid sequence is presented as SEQ ID NO. 95.

OspA-K48/OspA-B31/OspA-K48 A chimer of OspA from strain B31 (OspA-B31) and OspA from strain K48 (OspA-K48) was generated using the method described above. This chimeric nucleic acid sequence included bp 1-570 from OspA-B31, followed by bp 570-651 from OspA-B31, followed by bp 650-820 from OspA-K48. Primers used included: the fusion primer, 5'-CCCCAGATTTTGAAATCTTGCTTAAAACAAC-3' (SEQ ID NO.96); and the sequence primer, #357 (SEQ ID NO.15). The chimeric nucleic acid sequence is presented as SEQ ID NO.97; the chimeric protein encoded by this chimeric nucleic acid sequence is presented as SEQ ID NO.98.

OspA-B31/OspA-K48/OspA-B31/OspA-K48

from strain B31 (OspA-B31) and OspA from strain K48 (OspA-K48) was generated using the method described above. This chimeric nucleic acid sequence included bp 1-420 from OspA-B31, followed by 420-570 from OspA-K48, followed by bp 570-650 from OspA-B31, followed by bp 651-820 from OspA-K48. Primers used included: the fusion primer, 5'-CAAGTCTGGTTCCAATTTGCTCTTGTTATTAT-3' (minus strand primer #436-420) (SEQ ID NO. 99); and the sequence primer, #357 (SEQ ID NO. 15). The chimeric nucleic acid sequence is presented as SEQ ID NO. 100; the chimeric protein encoded by this chimeric nucleic acid sequence is presented as SEQ ID NO. 101.

OSPA-B31/OSPB-B31 A chimer of OSPA and OSPB from strain
B31 (OSPA-B31, OSPB-B31) was generated using the method
described above. The chimeric nucleic acid sequence
included bp 1-651 from OSPA-B31, followed by bp 652-820
from OSPB-B31. Primers used included: the fusion primer,

5'-GTTAAAGTGCTAGTACTGTCATTCCAAGCTGCAGTTTTTTT-3' (minus
strand primer #740-651) (SEQ ID NO. 102); the carboxyterminal sequence of OSPB primer #1106 (SEQ ID NO. 25); and
the sequence primer #357 (SEQ ID NO. 15). The chimeric

nucleic acid sequence is presented as SEQ ID NO. 103; the chimeric protein encoded by this chimeric nucleic acid sequence is presented as SEQ ID NO. 104.

OspA-B31/OspB-B31/OspC-B31 A chimer of OspA, OspB and
OspC from strain B31 (OspA-B31, OspB-B31, and OspC-B31) was
generated using the method described above. The chimeric
nucleic acid sequence included bp 1-650 from OspA-B31,
followed by bp 652-820 from OspB-B31, followed by bp 74-630
of OspC-B31. Primers used included: the fusion primer, 5'TGCAGATGTAATCCCATCCGCCATTTTTAAAGCGTTTTT-3' (SEQ ID NO.
105); and the carboxy-terminal sequence of OspC primer (SEQ
ID NO. 28). The chimeric nucleic acid sequence is
presented as SEQ ID NO. 106; the chimeric protein encoded
by this chimeric nucleic acid sequence is presented as SEQ
ID NO. 107.

OspC-B31/OspA-B31/OspB-B31 A chimer of OspA, OspB and OspC from strain B31 (OspA-B31, OspB-B31, and OspC-B31) was generated using the method described above. The chimeric nucleic acid sequence included bp 1-630 from OspC-B31, followed by bp 52-650 from OspA-B31, followed by bp 650-820 of OspB-B31. Primers used included: the amino-terminal sequence of OspC primer having SEQ ID NO. 27; the fusion primer, 5'-GCTGCTAACATTTTGCTTAGGTTTTTTTGGACTTTC-3' (minus strand primer #69-630) (SEQ ID NO. 108); and the sequence primers #520 (SEQ ID NO. 40) and #200 (SEQ ID NO. 18). The chimeric nucleic acid sequence is presented as SEQ ID NO. 109; the chimeric protein encoded by this chimeric nucleic acid sequence is presented as SEQ ID NO. 110.

30 Additional Chimeric Nucleic Acid Sequences

Using the methods described above, other chimeric nucleic acid sequences were produced. These chimeric

nucleic acid sequences, and the proteins encoded, are summarized in Table 3.

Table III Chimeric Nucleic acid Sequences and the Encoded Proteins

	1				
Chimers Generated (base pairs)	SEQ ID NO. (nt)	SEQ ID NO. (protein)			
OspA (52-882) / p93 (1168-2100)	111	112			
OspB (45-891) / p41 (122-234)	113	114			
OspB (45-891) / p41 (122-295)	115	116			
OspB (45-891) / p41 (140-234)	117	118			
OspB (45-891) / p41 (140-295)	119	120			
OspB (45-891) / p41 (122-234) / OspC (58-633)	121	122			
OspA-Tro/OspA-Bo	137	138			
OspA-PGau/OspA-Bo	139	140			
OspA-B31/OspA-PGau/OspA-B31/ OspA-K48	141	142			
OspA-PGau/OspA-B31/OspA-K48	143	144			

C. Purification of Proteins Generated by Chimeric Nucleic Acid Sequences

The chimeric nucleic acid sequences described above, as well as chimeric nucleic acid sequences produced by the methods described above, are used to produce chimeric proteins encoded by the nucleic acid sequences. Standard methods, such as those described above in Example 3, concerning the expression of proteins from Borrelia genes, can be used to express the proteins in a compatible host organism. The chimeric proteins can then be isolated and purified using standard techniques.

If the chimeric protein is soluble, it can be purified on a Sepharose column. Insoluble proteins can be solubilized in guanidine and purified on a Ni++ column;

alternatively, they can be solubilized in 10 mM NaPO₄ with 0.1 - 1% TRIXON X 114, and subsequently purified over an S column (Pharmacia). Lipidated proteins were generally purified by the latter method. Solubility was determined by separating both soluble and insoluble fractions of cell lysate on a 12% PAGE gel, and checking for the localization of the protein by Coomasie staining, or by Western blotting with monoclonal antibodies directed to an antigenic polypeptide of the chimeric protein.

10 Equivalents

Those skilled in the art will recognize, or be able to ascertain using no more than routine experimentation, many equivalents to the specific embodiments of the invention described herein. such equivalents are intended to be encompassed in the scope of the following claims.

-54-

CLAIMS

What is claimed is:

- A chimeric protein comprising two or more antigenic
 Borrelia polypeptides, wherein the antigenic Borrelia
 polypeptides which comprise the chimeric protein do
 not occur naturally in the same protein in Borrelia.
 - 2. The chimeric protein of Claim 1, wherein the antigenic Borrelia polypeptides are from two or more different species of Borrelia.
- The chimeric protein of Claim 2, wherein the antigenic Borrelia polypeptides are derived from Borrelia proteins selected from the group consisting of: outer surface protein A, outer surface protein B, outer surface protein C, outer surface protein D, p12, p39, p41, p66, and p93.
 - 4. The chimeric protein of Claim 3, wherein the antigenic Borrelia polypeptides are from corresponding proteins from two or more different species of Borrelia.
- 5. The chimeric protein of Claim 3, wherein the antigenic
 Borrelia polypeptides are from non-corresponding
 proteins from at least two different species of
 Borrelia.
 - 6. The chimeric protein of Claim 1, wherein two or more antigenic Borrelia polypeptides are from the same species of Borrelia.

PCT/US94/12352

- 7. The chimeric protein of Claim 6, wherein the antigenic Borrelia polypeptides are derived from Borrelia proteins selected from the group consisting of: outer surface protein A, outer surface protein B, outer surface protein C, outer surface protein D, pl2, p39, p41, p66, and p93.
 - 8. The chimeric protein of Claim 7, wherein the antigenic Borrelia polypeptides are from the same protein.
- 9. The chimeric protein of Claim 6, wherein the antigenic Borrelia polypeptides are from different proteins.
 - 10. A chimeric protein comprising two antigenic Borrelia polypeptides flanking a tryptophan residue, wherein the amino-proximal polypeptide consists of a polypeptide that is proximal from the single
- tryptophan residue of a first outer surface protein of Borrelia, and the carboxy-proximal polypeptide consists of a polypeptide that is distal from the single tryptophan residue of a second outer surface protein of Borrelia.
- 20 11. The chimeric protein of Claim 10, wherein the first and second outer surface proteins are from the same species of Borrelia.
- 12. The chimeric protein of Claim 11, wherein the first outer surface protein is outer surface protein A and the second outer surface protein is outer surface protein B.
 - 13. The chimeric protein of Claim 11, wherein the first outer surface protein is outer surface protein B, and

PCT/US94/12352

the second outer surface protein is outer surface protein A.

- 14. The chimeric protein of Claim 10, wherein the first and second outer surface proteins are from different species of *Borrelia*.
 - 15. The chimeric protein of Claim 14, wherein the first outer surface protein is outer surface protein A and the second outer surface protein is outer surface protein B.
- 10 16. The chimeric protein of Claim 14, wherein the first outer surface protein is outer surface protein B, and the second outer surface protein is outer surface protein A.
- 17. The chimeric protein of Claim 14, wherein the first
 and second outer surface proteins are corresponding
 proteins selected from the group consisting of: outer
 surface protein A and outer surface protein B.
- 18. The chimeric protein of Claim 10, wherein the first outer surface protein is outer surface protein A and the second outer surface protein is outer surface protein B.
- 19. The chimeric protein of Claim 18, wherein the aminoproximal polypeptide further comprises a first, second, and third hypervariable domain, the first hypervariable domain consisting of residues 120 through 140 of outer surface protein A, the second hypervariable domain consisting of residues 150 through 180 of outer surface protein A, and the third

hypervariable domain consisting of residues 200 through 217 of outer surface protein A.

- The chimeric protein of Claim 19, wherein the first and second hypervariable domains are derived from outer surface protein A from different species of Borrelia.
- The chimeric protein of Claim 10, further comprising an antigenic Borrelia polypeptide derived from a Borrelia protein selected from the group consisting of: outer surface protein A, outer surface protein B, outer surface protein C, outer surface protein D, p12, p39, p41, p66, and p93.
- 22. A nucleic acid sequence encoding a chimeric protein comprising two antigenic Borrelia polypeptides,
 wherein the two antigenic Borrelia polypeptides which comprise the chimeric protein do not occur naturally in the same protein in Borrelia.
- 23. The nucleic acid sequence of Claim 22, wherein the antigenic Borrelia polypeptides are from two or more different species of Borrelia.
- 24. The nucleic acid sequence of Claim 23, wherein the antigenic Borrelia polypeptides are derived from Borrelia proteins selected from the group consisting of: outer surface protein A, outer surface protein B, outer surface protein C, outer surface protein D, pl2, p39, p41, p66, and p93.
 - 25. The nucleic acid sequence of Claim 24, wherein the antigenic Borrelia polypeptides are from corresponding

proteins from two or more different species of Borrelia.

- 26. The nucleic acid sequence of Claim 24, wherein two or more of the antigenic *Borrelia* polypeptides are from non-corresponding proteins from different species of *Borrelia*.
- 27. The nucleic acid sequence of Claim 22, wherein two or more antigenic Borrelia polypeptides are from the same species of Borrelia.
- 10 28. The nucleic acid sequence of Claim 27, wherein the antigenic Borrelia polypeptides are derived from Borrelia proteins selected from the group consisting of: outer surface protein A, outer surface protein B, outer surface protein C, outer surface protein D, p12, p39, p41, p66, and p93.
 - 29. The nucleic acid sequence of Claim 28, wherein the antigenic *Borrelia* polypeptides are from the same protein.
- 30. The nucleic acid sequence of Claim 27, wherein the antigenic Borrelia polypeptides are from different proteins.

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- 31. A nucleic acid sequence encoding a chimeric protein comprising two antigenic Borrelia polypeptides flanking a tryptophan residue, wherein the aminoproximal polypeptide consists of a polypeptide that is proximal from the single tryptophan residue of a first outer surface protein of Borrelia, and the carboxy-proximal polypeptide consists of a polypeptide that is distal from the single tryptophan residue of a second outer surface protein of Borrelia.
- 10 32. The nucleic acid sequence of Claim 31, wherein the first and second outer surface proteins are from the same species of Borrelia.
- 33. The nucleic acid sequence of Claim 32, wherein the first outer surface protein is outer surface protein A and the second outer surface protein is outer surface protein B.
- 34. The nucleic acid sequence of Claim 32, wherein the first outer surface protein is outer surface protein B, and the second outer surface protein is outer surface protein A.
 - 35. The nucleic acid sequence of Claim 31, wherein the first and second outer surface proteins are from different species of Borrelia.
- 36. The nucleic acid sequence of Claim 35, wherein the
 first outer surface protein is outer surface protein A
 and the second outer surface protein is outer surface
 protein B.

WO 95/12676 PCT/US94/12352

-60-

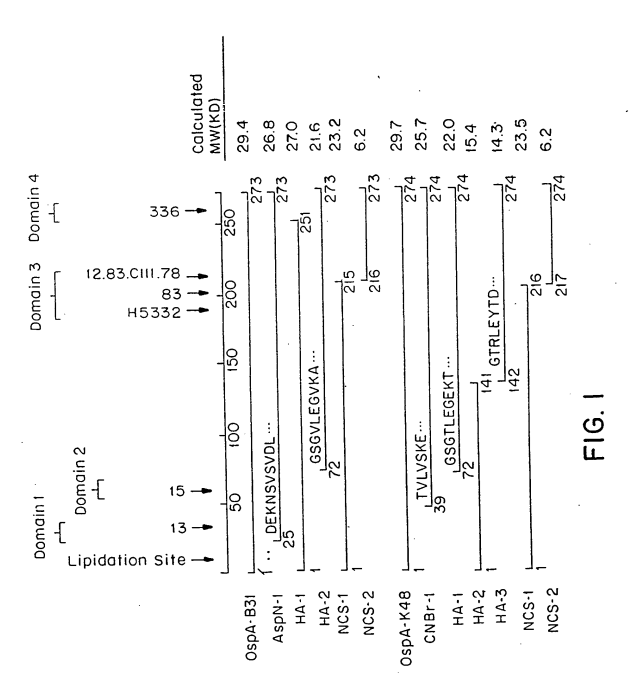
- 37. The nucleic acid sequence of Claim 35, wherein the first outer surface protein is outer surface protein B, and the second outer surface protein is outer surface protein A.
- 5 38. The nucleic acid sequence of Claim 35, wherein the first and second outer surface proteins are corresponding proteins selected from the group consisting of: outer surface protein A and outer surface protein B.
- 10 39. The nucleic acid sequence of Claim 31, wherein the first outer surface protein is outer surface protein A and the second outer surface protein is outer surface protein B.
- 40. The nucleic acid sequence of Claim 39, wherein the
 amino-proximal polypeptide further comprises a first
 and a second hypervariable domain, the first
 hypervariable domain consisting of amino acid residues
 1 through 140 of outer surface protein A, and the
 second hypervariable domain consisting of amino acid
 residues 150 through 217 of outer surface protein A.
 - 41. The nucleic acid sequence of Claim 40, wherein the first and second hypervariable domains are derived from outer surface protein A from different species of Borrelia.
- 25 42. The nucleic acid sequence of Claim 31, further comprising an antigenic Borrelia polypeptide derived from a Borrelia protein selected from the group consisting of: outer surface protein A, outer surface protein B, outer surface protein C, outer surface protein D, p12, p39, p41, p66, and p93.

- 43. A nucleic acid sequence having a sequence selected from the group consisting of: SEQ ID NO. 85, SEQ ID NO. 88, SEQ ID NO. 91, SEQ ID NO. 94, SEQ ID NO. 97, SEQ ID NO. 100, SEQ ID NO. 103, SEQ ID NO. 106, SEQ ID NO. 109, SEQ ID NO. 111, SEQ ID NO. 113, SEQ ID NO. 115, SEQ ID NO. 117, SEQ ID NO. 119, SEQ ID NO. 121, SEQ ID NO. 137, SEQ ID NO. 139, SEQ ID NO. 141, and SEQ ID NO. 143.
- 44. A protein having an amino acid sequence selected from the group consisting of: SEQ ID NO. 86, SEQ ID NO. 89, SEQ ID NO. 92, SEQ ID NO. 95, SEQ ID NO. 98, SEQ ID NO. 101, SEQ ID NO. 104, SEQ ID NO. 107, SEQ ID NO. 110, SEQ ID NO. 112, SEQ ID NO. 114, SEQ ID NO. 116, SEQ ID NO. 118, SEQ ID NO. 120, SEQ ID NO. 122, SEQ ID NO. 138, SEQ ID NO. 140, SEQ ID NO. 142, and SEQ ID NO. 144.
- 45. A chimeric protein according to any one of claims 1 to 21 and 44 for use in therapy or diagnosis, for example as a vaccine against <u>Borrelia</u> infection, in immunodiagnostic assays to detect the presence of antibodies to <u>Borrelia</u> or to measure T-cell reactivity.
- 46. A chimeric protein according to claim 45, wherein the immunodiagnostic assay is a dot blot, Western blot, ELISA or agglutination assay.

- 47. Use of the chimeric protein according to any one of claims 1 to 21 and 44, or the nucleic acid sequence of any one of claims 22 to 43, for the manufacture of a compound for use in therapy or diagnosis, for example as a vaccine against <u>Borrelia</u> infection, in immunodiagnostic assays to detect the presence of antibodies to <u>Borrelia</u> or to measure T-cell reactivity.
- 48. Use according to claim 47, wherein the

 immunodiagnostic assay is a dot blot, Western blot,

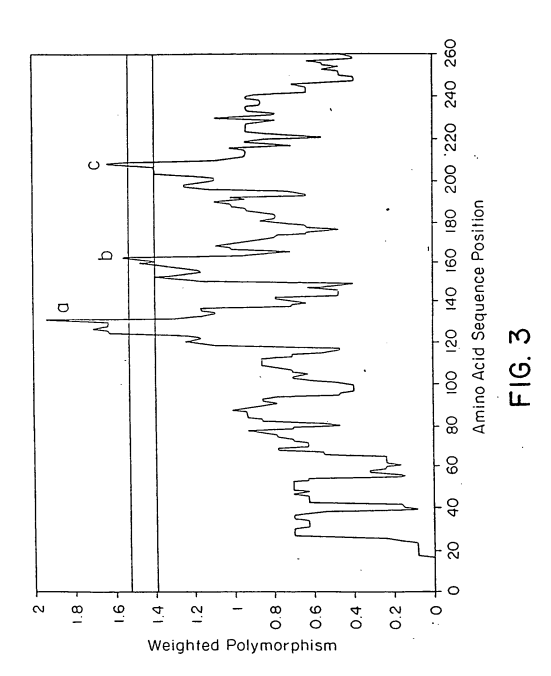
 ELISA or agglutination assay.



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	72	v	ဗ	ပ	ပ	ပ	ဖ	ဗ	v	Ö				,							
	1.1	z	z	z	z	z	z	z	z	z											
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Domain	69	×	×	~	- -	- -	~	~	~	~		270	KN	K	SAGINLEGKAVEITILKELKN	SAGTNLEGKAVEITTLKELKN	SAGTNLEGTAVEIKTLDELKN	SAGTNLEGTAVEIKTLDELKN	LLDELKN	SAGTNLEGKAVEITTLKELKN	SAGINLEGTAVEIKTLDEIKN
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	65 (-	_	-					TKLE	TNLE					SAGTNLEGTAVEIKTLDELKN		
	w	O	O	U	G	Ö	Ö	Ö	O	Ö		250	SNG	SAG							
		A-B31	A-TRO	A-K48	A-DK29	A-P/Gau	A-PKo	A-IP3	A-IP90	A-25015			A-B31	A-TRO	A-K48	A-DK29	A-P/Gau	A-PKo	A-IP3	A-IP90	A-25015
	41	ų	J	ı		ı	1	ı	1	1		220	ISGT	INS	SKT	SKT	EIAKSGEVTVALNDTNTTQATKKTGAWDSKT	SKT	SKT	SKT	AGT
	40	>	>	>	>	>	>	>	>	>			AAMN	GKWD	GKWD	NILKSGEITAALDDSDTTRATKKTGKWDSKT		GAWD	GAMD	GTWD	HISKSGEVTAELNDTDSTQATKKTGKWDAGT
	39	×	×	F	£	×	×	×	v	×		210	TKKT	TKKT	TKKT			TKKT	TKKT	TKKT	
in 1	38	Σ	Σ	Σ	Σ	Σ	Σ	н	Σ	Σ	m		SSAA	STQA	TTQA			TTOA	TTQA	TTQA	
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	36	v	Ö	ŋ	២	ບ່	v	უ	ဗ	Ö	Dom	200	NISKSGEVSVELNDTDSSAATKKTAAWNSGT	TVEL	TVAL			EIAKSGEVTVALNDTNTTQATKKTGAWDSKT	EIAKSGEVTVALNDTNTTQATKKTGAWDSKT	HISNSGEITVELNDSDTTQATKKTGTWDSKT	
	35	Qı	Q,	۵,	۵	ф	Д	Д	Д	Q			SGEV	HIPNSGEL	SGEI			SGEV	SGEV	SGEI	
	34	u	J	IJ	J	J	٦.	J	ı	H		190	NISK		NILK		EIAK	EIAK	EIAK	HISN	
		A-B31	A-TRO	A-K48	A-DK29	A-P/Gau	A-PKo	A-IP3	A-1P90	A-25015			A-B31	A-TRo	A-K48	A-DK29	A-P/Gau	A-PKo	A-IP3	A-1P90	A-25015

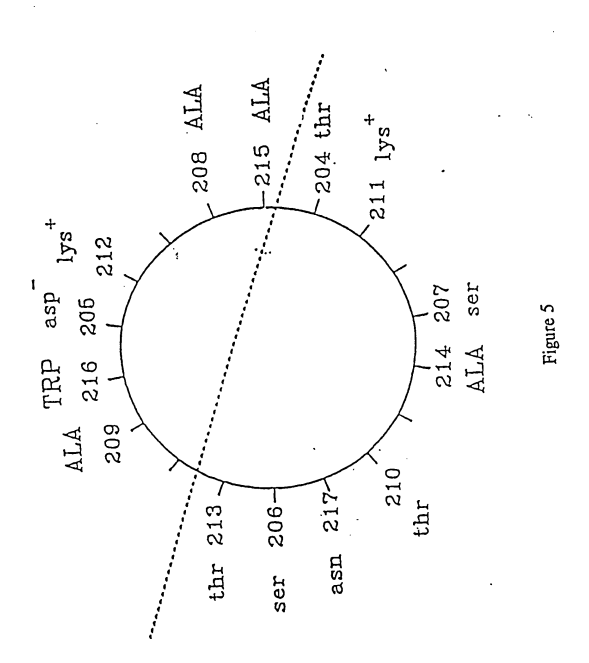
Figure 2



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B31:ELNDTDSSAATKKTAAWNSGTK48:ALDDSOTTQATKKTGKWDSKT613:ELNDSOTSAATKKTAAWNSGT625:ELNDTDSSAATKKTGKWNSGT640:ELNDTDSSAATKKTAAWDSKT613/625:ELNDSDISAATKKTGKWNSGT613/640:ELNDSOISAATKKTAAWDSKT

Figure 4



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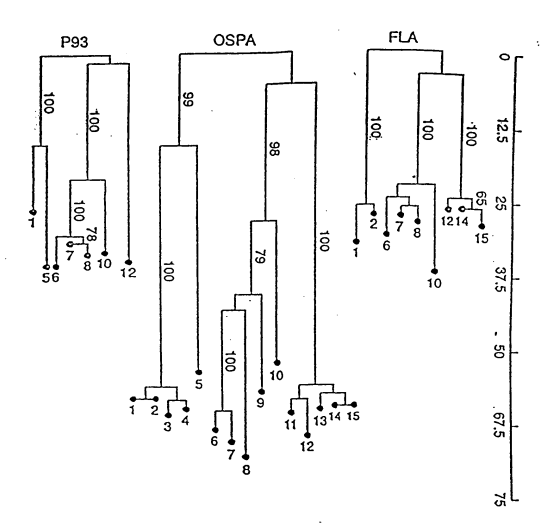


Figure 6

ATG Met 1	AAA Lys	AAA Lys	TAT Tyr	TTA Leu 5	TTG Leu	GGA Gly	ATA Ile	GGT Gly	CTA Leu 10	ATA Ile	TTA Leu	GCC Ala	TTA Leu	ATA Ile 15	GCA Ala		48
TGT Cys	AAG Lys	CAA Gln	AAT Asn 20	GTT Val	AGC Ser	AGC Ser	CTT Leu	GAC Asp 25	GAG Glu	AAA Lys	AAC Asn	AGC Ser	GTT Val 30	TCA Ser	GTA Val		96
GAT Asp	TTG Leu	CCT Pro 35	GGT Gly	GAA Glu	ATG Met	AAA Lys	GTT Val 40	CTT Leu	GTA Val	AGC Ser	Γλε, ΥΥΥ	GAA Glu 45	AAA Lys	AAC Asn	AAA Lys		144
G AC	GGC Gly 50	AAG Lys	TAC Tyr	GAT Asp	CTA Leu	ATT Ile 55	GCA Ala	ACA Thr	GTA Val	GAC Asp	AAG Lys 60	CTT Leu	GAG Glu	CTT Leu	TAE TAE	٠.	192
GGA Gly 65	ACT Thr	TCT Ser	GAT Asp	AAA Lys	AAC Asn 70	AAT Asn	GGA Gly	TCT Ser	GGA Gly	GTA Val 75	CTT Leu	GAA Glu	Gly	GTÀ Val	AAA Lys 80		240
GCT Ala	GAC Asp	AAA Lys	AGT Ser	AAA Lys 85	GTA Val	AAA Lys	TTA Leu	ACA Thr	ATT Ile 90	TCT Ser	GAC Asp	GAT Asp	CTA Leu	GGT Gly 95	CAA Gln		288
ACC Thr	ACA Thr	CTT Leu	GAA Glu 100	GTT Val	TTC Phe	AAA Lys	GAA Glu	GAT Asp 105	GGC Gly	Lys	ACA Thr	CTA Leu	GTA Val 110	TCA Ser	AAA Lys		336
AAA Lys	GTA Val	ACT Thr 115	TCC Ser	AAA Lys	GAC Asp	AAG Lys	TCA Ser 120	Ser	ACA Thr	GAA Glu	GAA Glu	AAA Lys 125	TTC Phe	AAT Asn	GAA Glu		384
AAA Lys	GGT Gly 130	Glu	GTA Val	TCT Ser	GAA Glu	AAA Lys 135	ATA Ile	ATA Ile	ACA Thr	AGA Arg	GCA Ala 140	Asp	GGA Gly	ACC Thr	AGA Arg		432
CTT Leu 145	Glu	TAC	ACA Thr	GGA Gly	ATT Ile 150	Lys	AGC Ser	GAT Asp	GGA Gly	TCT Ser 155	Gly	Lys Lys	GCT Ala	AAA Lys	GAG Glu 160		480
GTT Val	TTA Leu	AAA Lys	GGC Gly	TAT Tyr 165	Val	CTT Leu	GAA Glu	GGA Gly	ACT Thr 170	Leu	ACT Thr	GCT Ala	GAA Glu	AAA Lys 175	ACA Thr		528
ACA Thr	TTG Leu	GTC Val	GTT Val	. Lys	GAA Glu	GGA Gly	ACT Thr	GTI Val 185	Thr	TTA Leu	AGC Ser	AAA Lys	AAT Asn 190	116	TCA Ser		576
AAA Lys	TCI Ser	GGG Gly 199	/ Glu	GTI Val	TCA Ser	GTI Val	GAP Glu 200	ı Lev	TAA 7 Asr	GAC Asp	ACI Thr	GAC Asp 205	Ser	AGT Ser	GCT Ala		624

Figure 7 (1 of 2)

							Thr			67.
							ACA Thr	GAA Glu 240	٠.	72
							AAA Lys			761
							AAC Asn 270			816
AAA Lvs	TA						· :			822

Figure 7 (2 of 2)

OSPA K48

ATG AAA AAA TAT TTA TTG GGA ATA GGT CTA ATA TTA GCC TTA ATA GCA TAC TTT TTT ATA AAT AAC CCT TAT CCA GAT TAT AAT CGG AAT TAT CGT Met Lys Lys Tyr Leu Leu Gly Ile Gly Leu Ile Leu Ala Leu Ile Ala>

50 60 70 80 90

TGT AAG CAA AAT GTT AGC AGC CTT GAT GAA AAA AAT AGC GTT TCA GTA ACA TTC GTT TTA CAA TCG TCG GAA CTA CTT TTT TTA TCG CAA AGT CAT Cys Lys Gln Asn Val Ser Ser Leu Asp Glu Lys Asn Ser Val Ser Val>

100 110 120 130 140

GAT TTA CCT GGT GGA ATG ACA GTT CTT GTA AGT AAA GAA AAA GAC AAA CTA AAT GGA CCA CCT TAC TGT CAA GAA CAT TCA TTT CTT TTT CTG TTT Asp Leu Pro Gly Gly Met Thr Val Leu Val Ser Lys Glu Lys Asp Lys>

150 160 170 180 190

GAC GGT AAA TAC AGT CTA GAG GCA ACA GTA GAC AAG CTT GAG CTT AAA CTG CCA TTT ATG TCA GAT CTC CGT TGT CAT CTG TTC GAA CTC GAA TTT ASP Gly Lys Tyr Ser Leu Glu Ala Thr Val Asp Lys Leu Glu Leu Lys>

200 210 220 230 240

GGA ACT TCT GAT AAA AAC AAC GGT TCT GGA ACA CTT GAA GGT GAA AAA CCT TGA AGA CTA TTT TTG TTG CCA AGA CCT TGT GAA CTT CCA CTT TTT Gly Thr Ser Asp Lys Asn Asn Gly Ser Gly Thr Leu Glu Gly Glu Lys>

250 260 270 280

ACT GAC AAA AGT AAA GTA AAA TTA ACA ATT GCT GAT GAC CTA AGT CAA TGA CTG TTT TCA TTT CAT TTT AAT TGT TAA CGA CTA CTG GAT TCA GTT Thr Asp Lys Ser Lys Val Lys Leu Thr Ile Ala Asp Asp Leu Ser Gln>

290 300 310 320 330

ACT AAA TTT GAA ATT TTC AAA GAA GAT GCC AAA ACA TTA GTA TCA AAA TGA TTT AAA CTT TAA AAG TTT CTT CTA CGG TTT TGT AAT CAT AGT TTT Thr Lys Phe Glu Ile Phe Lys Glu Asp Ala Lys Thr Leu Val Ser Lys>

340 350 360 370 380

AAA GTA ACC CTT AAA GAC AAG TCA TCA ACA GAA GAA AAA TTC AAC GAA TTT CAT TGG GAA TTT CTG TTC AGT AGT TGT CTT CTT TTT AAG TTG CTT Lys Val Thr Leu Lys Asp Lys Ser Ser Thr Glu Glu Lys Phe Asn Glu>

FIGURE 8 (1 of 3)

WO 95/12676 PCT/US94/12352

10/133

OSP A K48

400 390 430 AAG GGT GAA ACA TCT GAA AAA ACA ATA GTA AGA GCA AAT GGA ACC AGA TTC CCA CTT TGT AGA CTT TTT TGT TAT CAT TCT CGT TTA CCT TGG TCT Lys Gly Glu Thr Ser Glu Lys Thr Ile Val Arg Ala Asn Gly Thr Arg> 460 440 450 470 480 CTT GAA TAC ACA GAC ATA AAA AGC GAT GGA TCC GGA AAA GCT AAA GAA GAA CTT ATG TGT CTG TAT TTT TCG CTA CCT AGG CCT TTT CGA TTT CTT Leu Glu Tyr Thr Asp Ile Lys Ser Asp Gly Ser Gly Lys Ala Lys Glu> 500 510 490 520 GTT TTA AAA GAC TTT ACT CTT GAA GGA ACT CTA GCT GCT GAC GGC AAA CAA AAT TTT CTG AAA TGA GAA CTT CCT TGA GAT CGA CGA CTG CCG TTT Val Leu Lys Asp Phe Thr Leu Glu Gly Thr Leu Ala Ala Asp Gly Lys> 530 540 550 560 570 ACA ACA TTG AAA GTT ACA GAA GGC ACT GTT GTT TTA AGC AAG AAC ATT TGT TGT AAC TTT CAA TGT CTT CCG TGA CAA CAA AAT TCG TTC TTG TAA Thr Thr Leu Lys Val Thr Glu Gly Thr Val Val Leu Ser Lys Asn Ile> 580 600 590 610 620 TTA AAA TCC GGA GAA ATA ACA GTT GCA CTT GAT GAC TCT GAC ACT ACT AAT TTT AGG CCT CTT TAT TGT CAA CGT GAA CTG CTG AGA CTG TGA TGA Leu Lys Ser Gly Glu Ile Thr Val Ala Leu Asp Asp Ser Asp Thr Thr> 650 630 640 660 670 CAG GCT ACT AAA AAA ACT GGA AAA TGG GAT TCA AAA ACT TCC ACT TTA GTC CGA TGA TTT TTT TGA CCT TTT ACC CTA AGT TTT TGA AGG TGA AAT Gln Ala Thr Lys Lys Thr Gly Lys Trp Asp Ser Lys Thr Ser Thr Leu> 700 680 690 710 720 ACA ATT AGT GTG AAT AGC CAA AAA ACC AAA AAC CTT GTA TTC ACA AAA TGT TAA TCA CAC TTA TCG GTT TTT TGG TTT TTG GAA CAT AAG TGT TTT Thr Ile Ser Val Asn Ser Gln Lys Thr Lys Asn Leu Val Phe Thr Lys> 730 740 750 760 GAA GAC ACA ATA ACA GTA CAA AAA TAC GAC TCA GCA GGC ACC AAT CTA CTT CTG TGT TAT TGT CAT GTT TTT ATG CTG AGT CGT CCG TGG TTA GAT

FIGURE 8 (2 of 3)

Glu Asp Thr Ile Thr Val Gln Lys Tyr Asp Ser Ala Gly Thr Asn Leu>

Osp A K-48

770 780 790 800 810

GAA GGC AAA GCA GTC GAA ATT ACA ACA CTT AAA GAA CTT AAA AAC GCT

GAA TTT CTT GAA TTT TTG CGA

CTT CCG TTT CGT CAG CTT TAA TGT TGT GAA TTT CTT GAA TTT TTG CGA Glu Gly Lys Ala Val Glu Ile Thr Thr Leu Lys Glu Leu Lys Asn Ala>

OSP A K48

. .

820

TTA AAA TAA AAT TTT ATT Leu Lys ***>

FIGURE 8 (3 of 3)

OSP A PGAU

10	20 . 30	40
ATG AAA AAA TAT TTA TTG		
TAC TTT TTT ATA AAT AAC		TAT AAT CGG AAT TAT CGT Ile Leu Ala Leu Ile Ala>
met bys bys Tyl bea bea	. 01, 110 01, 100	and but the sea lie May
50 60	70	80 90
TGC AAG CAA AAT GTT AGC		
ACG TTC GTT TTA CAA TCG		
Cys Lys Gln Asn Val Ser	Ser Leu Asp Giu	Lys Asn Ser Ala Ser Val>
100 110	120	130 140
	• • • •	* * * *
GAT TTG CCT GGT GAG ATG CTA AAC GGA CCA CTC TAC		
		Ser Lys Glu Lys Asp Lys>
150 160	170	180 190
	•	•
GAC GGT AAG TAC AGT CTA		
CTG CCA TTC ATG TCA GAT		Asp Lys Ile Glu Leu Lys>
Asp Gly Lys Tyr Ser Led	1 Lys Ale III. Val	was the did fed five
200 210	220	230 240
GGA ACT TCT GAT AAA GAC	AAT GGT TCT GGA	GTG CTT GAA GGT ACA AAA
CCT TGA AGA CTA TTT CTG	TTA CCA AGA CCT	CAC GAA CTT CCA TGT TTT
Gly Thr Ser Asp Lys Asp	Asn Gly Ser Gly	Val Leu Glu Gly Thr Lys>
250	260 270	· 280
Chr ChC AAA ACT AAA GC2	דדג גוג גדד מממ מ	GCT GAC GAT CTA AGT AAA
CTA CTG TTT TCA TTT CG1		
		CON CIG CIN ONL ICK III
Wah wah raa ser raa wre	a Lys Leu Thr Ile	Ala Asp Asp Leu Ser Lys>
290 300		
290 300	310	Ala Asp Asp Leu Ser Lys> 320 330
290 300 ACC ACA TTC GAA CTT TTA	310	Ala Asp Asp Leu Ser Lyss 320 330 AAA ACA TTA GTG TCA AGA
290 300 ACC ACA TTC GAA CTT TTA TGG TGT AAG CTT GAA AAA	310	Ala Asp Asp Leu Ser Lys> 320 330
290 300 ACC ACA TTC GAA CTT TTA TGG TGT AAG CTT GAA AAT Thr Thr Phe Glu Leu Leu	310	Ala Asp Asp Leu Ser Lyss 320 330 AAA ACA TTA GTG TCA AGA TTT TGT AAT CAC AGT TCT
290 300 ACC ACA TTC GAA CTT TTA TGG TGT AAG CTT GAA AAA Thr Thr Phe Glu Leu Leu 340 350	310 3 A AAA GAA GAT GGC T TTT CTT CTA CCG u Lys Glu Asp Gly 360	Ala Asp Asp Leu Ser Lyss 320 330 AAA ACA TTA GTG TCA AGA TTT TGT AAT CAC AGT TCT Lys Thr Leu Val Ser Args 370 380
290 300 ACC ACA TTC GAA CTT TTX TGG TGT AAG CTT GAA AAX Thr Thr Phe Glu Leu Leu 340 350 AAA GTA AGT TCT AGA GAG	310 A AAA GAA GAT GGC T TTT CTT CTA CCG U Lys Glu Asp Gly 360 C AAA ACA TCA ACA	Ala Asp Asp Leu Ser Lyss 320 330 AAA ACA TTA GTG TCA AGA TTT TGT AAT CAC AGT TCT Lys Thr Leu Val Ser Args 370 380 GAT GAA ATG TTC AAT GAA
290 300 ACC ACA TTC GAA CTT TTA TGG TGT AAG CTT GAA AAA Thr Thr Phe Glu Leu Leu 340 350 AAA GTA AGT TCT AGA GAA TTT CAT TCA AGA TCT CTO	310 A AAA GAA GAT GGC T TTT CTT CTA CCG U Lys Glu Asp Gly 360 C AAA ACA TCA ACA G TTT TGT AGT TGT	Ala Asp Asp Leu Ser Lyss 320 330 AAA ACA TTA GTG TCA AGA TTT TGT AAT CAC AGT TCT Lys Thr Leu Val Ser Args 370 380

FIGURE 9 (1 of 3)

430

OSP A P

400 390 AAA GGT GAA TTG TCT GCA AAA ACC ATG ACA AGA GAA AAT GGA ACC AAA TTT CCA CTT AAC AGA CGT TTT TGG TAC TGT TCT CTT TTA CCT TGG TTT Lys Gly Glu Leu Ser Ala Lys Thr Het Thr Arg Glu Asn Gly Thr Lys> 470 480 460 450 440 CTT GAA TAT ACA GAA ATG AAA AGC GAT GGA ACC GGA AAA GCT AAA GAA GAA CTT ATA TGT CTT TAC TTT TCG CTA CCT TGG CCT TTT CGA TTT CTT Leu Glu Tyr Thr Glu Met Lys Ser Asp Gly Thr Gly Lys Ala Lys Glu> 510 520 500 490 GTT TTA AAA AAG TTT ACT CTT GAA GGA AAA GTA GCT AAT GAT AAA GTA CAA AAT TIT TIC AAA TGA GAA CIT CCT TIT CAT CGA TIA CIA TIT CAT Val Leu Lys Lys Phe Thr Leu Glu Gly Lys Val Ala Asn Asp Lys Val> 570 560 550 530 540 ACA TTG GAA GTA AAA GAA GGA ACC GTT ACT TTA AGT AAG GAA ATT GCA TGT AAC CTT CAT TTT CTT CCT TGG CAA TGA AAT TCA TTC CTT TAA CGT Thr Leu Glu Val Lys Glu Gly Thr Val Thr Leu Ser Lys Glu Ile Ala> 620 610 600 590 AAA TCT GGA GAA GTA ACA GTT GCT CTT AAT GAC ACT AAC ACT ACT CAG TTT AGA CCT CTT CAT TGT CAA CGA GAA TTA CTG TGA TTG TGA TGA GTC Lys Ser Gly Glu Val Thr Val Ala Leu Asn Asp Thr Asn Thr Thr Gln> 660 650 640 630 GCT ACT AAA AAA ACT GGC GCA TGG GAT TCA AAA ACT TCT ACT TTA ACA CGA TGA TTT TTT TGA CCG CGT ACC CTA AGT TTT TGA AGA TGA AAT TGT Ala Thr Lys Lys Thr Gly Ala Trp Asp Ser Lys Thr Ser Thr Leu Thr> 720 700 710 680 690 ATT AGT GTT AAC AGC AAA AAA ACT ACA CAA CTT GTG TTT ACT AAA CAA TAA TCA CAA TTG TCG TTT TTT TGA TGT GTT GAA CAC AAA TGA TTT GTT Ile Ser Val Asn Ser Lys Lys Thr Thr Gln Leu Val Phe Thr Lys Gln> 750 760 740 730 TAC ACA ATA ACT GTA AAA CAA TAC GAC TCC GCA GGT ACC AAT TTA GAA ATG TGT TAT TGA CAT TTT GTT ATG CTG AGG CGT CCA TGG TTA AAT CTT

Tyr Thr Ile Thr Val Lys Gln Tyr Asp Ser Ala Gly Thr Asn Leu Glu>

WO 95/12676 PCT/US94/12352

14/133

OSP A PGAU

770 780 790 800 810

GGC ACA GCA GTC GAA ATT AAA ACA CTT GAT GAA CTT AAA AAC GCT TTA CCG TGT CGT CAG CTT TAA TTT TGT GAA CTA CTT GAA TTT TTG CGA AAT Gly Thr Ala Val Glu Ile Lys Thr Leu Asp Glu Leu Lys Asn Ala Leu>

820

AAA TAA TTT ATT Lys ***>

FIGURE 9 (3 of 3)

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ATG Met 1	AAA Lys	AAA Lys	TAT Tyr	TTA Leu 5	TTG Leu	GGA Gly	ATA Ile	GGT Gly	CTA Leu 10	ATA Ile	TTA Leu	GCT Ala	TTA Leu	ATA Ile 15	GCA Ala	48
Cys	AAG Lys	CAA Gln	AAT Asn 20	GTT Val	AGC Ser	AGC Ser	CTT Leu	GAC Asp 25	GAG Glu	AAA Lys	AAC Asn	AGC Ser	GTT Val 30	TCA Ser	GTA Val	96
GAT Asp	TTG Leu	CCT Pro 35	GGT Gly	GAA Glu	ATG Met	AAA Lys	GTT Val 40	CTT Leu	GTA Val	AGC Ser	AAA Lys	GAA Glu 45	AAA Lys	GAC Asp	AAA Lys	144
GAC Asp	GGC Gly 50	AAG Lys	TAC Tyr	AGT Ser	CTA Leu	ATG Met 55	GCA Ala	ACA Thr	GTA Val	GAC Asp	AAG Lys 60	CTT	GAG Glu	CTT Leu	AAA Lys	192

Figure 10 (1 of 2)

GGA Gly 65	ACA Thr	TCT Ser	GAT Asp	AAA Lys	AAC Asn 70	TAA neA	GGA Gly	TCT Ser	GGG Gly	GTG Val 75	CTT Leu	GAA Glu	GGC Gly	GTA Val	AAA Lys 80		240
GCT Ala	GAC Asp	AAA Lys	AGC Ser	AAA Lys 85	GTA Val	AAA Lys	TTA Leu	ACA Thr	GTT Val 90	TCT Ser	Asp	GAT Asp	CTA Leu	AGC Ser 95	ACA Thr		288
ACC Thr	ACA Thr	CTT Leu	GAA Glu 100	GTT Val	TTA Leu	AAA Lys	GAA Glu	GAT Asp 105	GGC Gly	TÀ8 TÀ8	ACA Thr	TTA Leu	GTG Val 110	TCA Ser	AAA Lys		336
AAA Lys	AGA Arg	ACT Thr 115	TCT Ser	AAA Lys	GAT Așp	AAG Lys	TCA Ser 120	TCA Ser	ACA Thr	GAA Glu	GAA Glu	AAG Lys 125	TTC Phe	AAT Asn	GĀA Glu	•.	384
AAA Lys	GGC Gly 130	GAA Glu	TTA Leu	GTT Val	GAA Glu	AAA Lys 135	ATA Ile	ATG Met	GCA Ala	AGA Arg	GCA Ala 140	AAC Asn	GGA Gly	ACC Thr	ATA Ile		432
CTT Leu 145	GAA Glu	TAC Tyr	ACA Thr	GGA Gly	ATT Ile	AAA Lys	AGC Ser	GAT Asp	GGA Gly	TCC Ser 155	GGA Gly	AAA Lys	GCT Ala	AAA Lys	GAA Glu 160		480
ACT	TTA Leu	TAa YYY	GAA Glu	TAT Tyr 165	GTT Val	CTT Leu	GAA Glu	GGA Gly	ACT Thr 170	CTA Leu	ACT Thr	GCT Ala	GAA Glu	AAA Lys 175	GCA Ala		528
ACA Thr	TTG Leu	GTG Val	GTT Val 180	Ь Гув	GAA Glu	GGA Gly	ACT Thr	GTT Val 185	ACT Thr	TTA Leu	AGT Ser	AAG Lys	CAC His 190	ATT Ile	TCA Ser		576
AAA Lys	TCT Ser	GGA Gly 195	GAA Glu	GTA Val	ACA Thr	GCT Ala	GAA Glu 200	CTT Leu	AAT Asn	GAC Asp	ACT Thr	GAC Asp 205	AGT Ser	ACT Thr	CAA Gln		624
GCT Ala	ACT Thr 210	Lys	AAA Lys	ACT Thr	GGG Gly	AAA Lys 215	TGG Trp	GAT Asp	GCA Ala	GGC Gly	ACT Thr 220	TCA Ser	ACT Thr	TTA Leu	ACA Thr		672
ATT Ile 225	Thr	GTA Val	AAC	AAC Asn	AAA Lys 230	Lys	ACT Thr	AAA Lys	GCC Ala	CTT Leu 235	Val	TTT Phe	ACA Thr	AAA Lys	CAA Gln 240		720
GAC Asp	ACA Thr	ATT Ile	ACA Thr	TCA Ser 245	Gln	AAA Lys	TAC Tyr	GAC	TCA Ser 250	Ala	GGA Gly	ACC	AAC Asn	TTG Leu 255	GAA Glu		768
GGC Gly	ACA	GCA Ala	GTC Val 260	Glu	ATT	AAA Lys	ACA Thr	CTT Leu 265	Asp	GAA Glu	CTT	AAA Lys	AAC Asn 270	Ala	TTA Leu		816
AGA Arg						-											819

Figure 10 (2 of 2)

^!!!!\TTT!!\T A!!=== /=... = ^-:

17//33

30 20 10 ATG AGA TTA TTA ATA GGA TTT GCT TTA GCG TTA GCT TTA ATA GGA TGT TAC TOT AAT AAT TAT COT AAA CGA AAT CGC AAT CGA AAT TAT COT ACA Met Arg Leu Leu Ile Gly Phe Ala Leu Ala Leu Ala Leu Ile Gly Cys> 90 80 70 60 GCA CAA AAA GGT GCT GAG TCA ATT GGT TCT CAA AAA GAA AAT GAT CTA CGT GTT TTT CCA CGA CTC AGT TAA CCA AGA GTT TTT CTT TTA CTA GAT Ala Gln Lys Gly Ala Glu Ser Ile Gly Ser Gln Lys Glu Asn Asp Leu> 140 130 120 110 AAC CTT GAA GAC TCT AGT AAA AAA TCA CAT CAA AAC GCT AAA CAA GAC TTG GAA CTT CTG AGA TCA TTT TTT AGT GTA GTT TTG CGA TTT GTT CTG Asn Leu Glu Asp Ser Ser Lys Lys Ser His Gln Asn Ala Lys Gln Asp> 170 190 160 CTT CCT GCG GTG ACA GAA GAC TCA GTG TCT TTG TTT AAT GGT AAT AAA GAA GGA CGC CAC TGT CTT CTG AGT CAC AGA AAC AAA TTA CCA TTA TTT Leu Pro Ala Val Thr Glu Asp Ser Val Ser Leu Phe Asm Gly Asm Lys> 230 220 210 200 • ATT TIT GTA AGC AAA GAA AAA AAT AGC TCC GGC AAA TAT GAT TTA AGA TAA AAA CAT TCG TTT CTT TTT TTA TCG AGG CCG TTT ATA CTA AAT TCT Ile Phe Val Ser Lys Glu Lys Asn Ser Ser Gly Lys Tyr Asp Leu Arg> 290 270 260 250 GCA ACA ATT GAT CAG GTT GAA CTT AAA GGA ACT TCC GAT AAA AAC AAT CGT TGT TAA CTA GTC CAA CTT GAA TTT CCT TGA AGG CTA TTT TTG TTA Ala Thr Ile Asp Gln Val Glu Leu Lys Gly Thr Ser Asp Lys Asn Asn> 320 310 300 290 GGT TCT GGA ACC CTT GAA GGT TCA AAG CCT GAC AAG AGT AAA GTA AAA CCA AGA CCT TGG GAA CTT CCA AGT TTC GGA CTG TTC TCA TTT CAT TTT Gly Ser Gly Thr Leu Glu Gly Ser Lys Pro Asp Lys Ser Lys Val Lys>

TTA ACA GTT TCT GCT GAT TTA AAC ACA GTA ACC TTA GAA GCA TTT GAT AAT TGT CAA AGA CGA CTA AAT TTG TGT CAT TGG AAT CTT CGT AAA CTA Leu Thr Val Ser Ala Asp Leu Asn Thr Val Thr Leu Glu Ala Phe Asp> 390 400 410 420 430

360

350

340

370

GCC AGC AAC CAA AAA ATT TCA AGT AAA GTT ACT AAA AAA CAG GGG TCA CGG TCG TTG GTT TTT TAA AGT TCA TTT CAA TGA TTT TTT GTC CCC AGT Ala Ser Asn Gln Lys Ile Ser Ser Lys Val Thr Lys Lys Gln Gly Ser> 450 470 450 460 ATA ACA GAG GAA ACT CTC AAA GCT AAT AAA TTA GAC TCA AAG AAA TTA TAT TGT CTC CTT TGA GAG TTT CGA TTA TTT AAT CTG AGT TTC TTT AAT Ile Thr Glu Glu Thr Leu Lys Ala Asn Lys Leu Asp Ser Lys Lys Leu> 500 510 490 · · · ._• __ ACA AGA TCA AAC GGA ACT ACA CTT GAA TAC TCA CAA ATA ACA GAT GCT TGT TCT AGT TTG CCT TGA TGT GAA CTT ATG AGT GTT TAT TGT CTA CGA Thr Arg Ser Asn Gly Thr Thr Leu Glu Tyr Ser Gln Ile Thr Asp Ala> 5€0 570 550 540 • • • . GAC AAT GCT ACA AAA GCA GTA GAA ACT CTA AAA AAT AGC ATT AAG CTT CTG TTA CGA TGT TTT CGT CAT CTT TGA GAT TTT TTA TCG TAA TTC GAA Asp Asn Ala Thr Lys Ala Val Glu Thr Leu Lys Asn Ser Ile Lys Leu> 600 610 590 580 GAA GGA AGT CTT GTA GTC GGA AAA ACA ACA GTG GAA ATT AAA GAA GGT CTT CCT TCA GAA CAT CAG CCT TTT TGT TGT CAC CTT TAA TTT CTT CCA Glu Gly Ser Leu Val Val Gly Lys Thr Thr Val Glu Ile Lys Glu Gly> 640 650 660 ACT GTT ACT CTA AAA AGA GAA ATT GAA AAA GAT GGA AAA GTA AAA GTC TGA CAA TGA GAT TTT TCT CTT TAA CTT TTT CTA CCT TTT CAT TTT CAG Thr Val Thr Leu Lys Arg Glu Ile Glu Lys Asp Gly Lys Val Lys Val> 690 700 710 TTT TTG AAT GAC ACT GCA GGT TCT AAC AAA AAA ACA GGT AAA TGG GAA AAA AAC TTA CTG TGA CGT CCA AGA TTG TTT TTT TGT CCA TTT ACC CTT Phe Leu Asn Asp Thr Ala Gly Ser Asn Lys Lys Thr Gly Lys Trp Glu> 750 760 740 GAC AGT ACT AGC ACT TTA ACA ATT AGT GCT GAC AGC AAA AAA ACT AAA CTG TCA TGA TCG TGA AAT TGT TAA TCA CGA CTG TCG TTT TTT TGA TTT Asp Ser Thr Ser Thr Leu Thr Ile Ser Ala Asp Ser Lys Lys Thr Lys> 790 800 810 780 GAT TTG GTG TTC TTA ACA GAT GGT ACA ATT ACA GTA CAA CAA TAC AAC CTA AAC CAC AAG AAT TGT CTA CCA TGT TAA TGT CAT GTT ATG TTG

Asp Leu Val Phe Leu Thr Asp Gly Thr Ile Thr Val Gln Gln Tyr Asn>

19/133

820 830 840 850 860

ACA GCT GGA ACC AGC CTA GAA GGA TCA GCA AGT GAA ATT AAA AAT CTT TGT CGA CCT TGG TCG GAT CTT CCT AGT CGT TCA CTT TAA TTT TTA GAA Thr Ala Gly Thr Ser Leu Glu Gly Ser Ala Ser Glu Ile Lys Asn Leu>

870 880 890

TCA GAG CTT AAA AAC GCT TTA AAA TAA AGT CTC GAA TTT TTG CGA AAT TTT ATT Ser Glu Leu Lys Asn Ala Leu Lys ***>

FIGURE 11 (3 of 3)

OspC-B31

Sequence Range: 1 to 633

10 20 30 40

ATG AAA AAG AAT ACA TTA AGT GCG ATA TTA ATG ACT TTA TTT TTA TTT TAC TTT TTC TTA TGT AAT TCA CGC TAT AAT TAC TGA AAT AAA AAT AAA Met Lys Lys Asn Thr Leu Ser Ala Ile Leu Met Thr Leu Phe Leu Phe>

50 60 70 80 90

ATA TCT TGT AAT AAT TCA GGG AAA GAT GGG AAT ACA TCT GCA AAT TCT TAT AGA ACA TTA TTA AGT CCC TTT CTA CCC TTA TGT AGA CGT TTA AGA Ile Ser Cys Asn Asn Ser Gly Lys Asp Gly Asn Thr Ser Ala Asn Ser>

100 110 120 130 140

GCT GAT GAG TCT GTT AAA GGG CCT AAT CTT ACA GAA ATA AGT AAA AAA CGA CTA CTC AGA CAA TTT CCC GGA TTA GAA TGT CTT TAT TCA TTT TTT Ala Asp Glu Ser Val Lys Gly Pro Asn Leu Thr Glu Ile Ser Lys Lys>

150 160 170 180 190

ATT ACG GAT TCT AAT GCG GTT TTA CTT GCT GTG AAA GAG GTT GAA GCG TAA TGC CTA AGA TTA CGC CAA AAT GAA CGA CAC TTT CTC CAA CTT CGC Ile Thr Asp Ser Asn Ala Val Leu Leu Ala Val Lys Glu Val Glu Ala>

200 210 220 230 240

TTG CTG TCA TCT ATA GAT GAA ATT GCT GCT AAA GCT ATT GGT AAA AAA AAC GAC AGT AGA TAT CTA CTT TAA CGA CGA TTT CGA TAA CCA TTT TTT Leu Leu Ser Ser Ile Asp Glu Ile Ala Ala Lys Ala Ile Gly Lys Lys>

250 260 270 280

ATA CAC CAA AAT AAT GGT TTG GAT ACC GAA TAT AAT CAC AAT GGA TCA TAT GTG GTT TTA TTA CCA AAC CTA TGG CTT ATA TTA GTG TTA CCT AGT Ile His Gln Asn Asn Gly Leu Asp Thr Glu Tyr Asn His Asn Gly Ser>

290 300 310 320 330

TTG TTA GCG GGA CGT TAT GCA ATA TCA ACC CTA ATA AAA CAA AAA TTA AAC AAT CGC CCT GCA ATA CGT TAT AGT TGG GAT TAT TTT GTT TTT AAT Leu Leu Ala Gly Arg Tyr Ala Ile Ser Thr Leu Ile Lys Gln Lys Leu>

340 350 360 370 380

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21/133

OBPC-B31

Lys Pro ***>

420 430 400 390 AAA TGT TCT GAA ACA TTT ACT AAT AAA TTA AAA GAA AAA CAC ACA GAT TTT ACA AGA CTT TGT AAA TGA TTA TTT AAT TTT CTT TTT GTG TGT CTA Lys Cys Ser Glu Thr Phe Thr Asn Lys Leu Lys Glu Lys His Thr Asp> 480 460 470 450 CTT GGT AAA GAA GGT GTT ACT GAT GCT GAT GCA AAA GAA GCC ATT TTA GAA CCA TTT CTT CCA CAA TGA CTA CGA CTA CGT TTT CTT CGG TAA AAT Leu Gly Lys Glu Gly Val Thr Asp Ala Asp Ala Lys Glu Ala Ile Leu> 510 --- . 500 490 AAA ACA AAT GGT ACT AAA ACT AAA GGT GCT GAA GAA CTT GGA AAA TTA TTT TGT TTA CCA TGA TTT TGA TTT CCA CGA CTT CTT GAA CCT TTT AAT Lys Thr Asn Gly Thr Lys Thr Lys Gly Ala Glu Glu Leu Gly Lys Leu> 560 550 540 530 TTT GAA TCA GTA GAG GTC TTG TCA AAA GCA GCT AAA GAG ATG CTT GCT AAA CTT AGT CAT CTC CAG AAC AGT TTT CGT CGA TTT CTC TAC GAA CGA Phe Glu Ser Val Glu Val Leu Ser Lys Ala Ala Lys Glu Met Leu Ala> 620 600 610 590 580 AAT TCA GTT AAA GAG CTT ACA AGC CCT GTT GTG GCA GAA AGT CCA AAA TTA AGT CAA TTT CTC GAA TGT TCG GGA CAA CAC CGT CTT TCA GGT TTT Asn Ser Val Lys Glu Leu Thr Ser Pro Val Val Ala Glu Ser Pro Lys> 630 AAA CCT TAA TTT GGA ATT

FIGURE 12 (2 of 2)

6

Osp 18 Sequence Range: 1 to 630



20 ATG AAA AAG AAT ACA TTA AGT GCG ATA TTA ATG ACT TTA TTT TTA TTT TAC TIT TTC TTA TGT AAT TCA CGC TAT AAT TAC TGA AAT AAA AAT AAA Met Lys Lys Asn Thr Leu Ser Ala Ile Leu Met Thr Leu Phe Leu Phe> 50 60 70 90 80 ATA TCT TGT AAT AAT TCA GGT GGG GAT ACC GCA TCT ACT AAT CCT GAT TAT AGA ACA TTA TTA AGT CCA CCC CTA TGG CGT AGA TGA TTA GGA CTA Ile Ser Cys Asn Asn Ser Gly Gly Asp Thr Ala Ser Thr Asn Pro Asp> 110 120 GAG TCT GCA AAA GGA CCT AAT CTT ACA GTA ATA AGC AAA AAA ATT ACA CTC AGA CGT TTT CCT GGA TTA GAA TGT CAT TAT TCG TTT TTT TAA TGT Glu Ser Ala Lys Gly Pro Asn Leu Thr Val Ile Ser Lys Lys Ile Thr> 160 150 170 180 190 GAT TCT AAT GCA TTT GTA CTG GCT GTG AAA GAA GTT GAG GCT TTG ATC CTA AGA TTA CGT AAA CAT GAC CGA CAC TTT CTT CAA CTC CGA AAC TAG Asp Ser Asn Ala Phe Val Leu Ala Val Lys Glu Val Glu Ala Leu Ile> 210 240 TCA TCT ATA GAT GAA CTT GCT AAT AAA GCT ATT GGT AAA GTA ATA CAT AGT AGA TAT CTA CTT GAA CGA TTA TTT CGA TAA CCA TTT CAT TAT GTA Ser Ser Ile Asp Glu Leu Ala Asn Lys Ala Ile Gly Lys Val Ile His> 250 260 270 CAA AAT AAT GGT TTA AAT GCT AAT GCG GGT CAA AAC GGA TCA TTG TTA GTT TTA TTA CCA AAT TTA CGA TTA CGC CCA GTT TTG CCT AGT AAC AAT Gln Asn Asn Gly Leu Asn Ala Asn Ala Gly Gln Asn Gly Ser Leu Leu> 290 300 310 320 330 GCA GGA GCC TAT GCA ATA TCA ACC CTA ATA ACA GAA AAA TTA AGT AAA CGT CCT CGG ATA CGT TAT AGT TGG GAT TAT TGT CTT TTT AAT TCA TTT Ala Gly Ala Tyr Ala Ile Ser Thr Leu Ile Thr Glu Lys Leu Ser Lys> 350 360 370 TTG AAA AAT TCA GAA GAG TTA AAT AAA AAA ATT GAA GAG GCT AAG AAC AAC TIT TTA AGT CTT CTC AAT TTA TTT TTT TAA CTT CTC CGA TTC TTG Leu Lys Asn Ser Glu Glu Leu Asn Lys Lys Ile Glu Glu Ala Lys Asn>

FIGURE 13 (1 of 2)

GGA ATT
Pro ***>

OspC-X48

23/133

410 420 390 400 CAT TCT GAA GCA TTT ACT AAT AGA CTA AAA GGT TCT CAT GCA CAA CTT GTA AGA CTT CGT AAA TGA TTA TCT GAT TTT CCA AGA GTA CGT GTT GAA His Ser Glu Ala Phe Thr Asn Arg Leu Lys Gly Ser His Ala Gln Leu> 450 460 440 470 GGA GTT GCT GCT ACT GAT GAT CAT GCA AAA GAA GCT ATT TTA AAG CCT CAA CGA CGA CGA TGA CTA CTA GTA CGT TTT CTT CGA TAA AAT TTC Gly Val Ala Ala Ala Thr Asp Asp His Ala Lys Glu Ala Ile Leu Lys> 490 500 ៊ី_{*} =:::: TCA AAT CCT ACT AAA GAT AAG GGT GCT AAA GCA CTT AAA GAC TTA TCT AGT TTA GGA TGA TTT CTA TTC CCA CGA TTT CGT GAA TTT CTG AAT AGA Ser Asn Pro Thr Lys Asp Lys Gly Ala Lys Ala Leu Lys Asp Leu Ser> 530 540 550 560 570 GAA TCA GTA GAA AGC TTG GCA AAA GCA GCG CAA GAA GCA TTA GCT AAT CTT AGT CAT CTT TCG AAC CGT TTT CGT CGC GTT CTT CGT AAT CGA TTA Glu Ser Val Glu Ser Leu Ala Lys Ala Ala Gln Glu Ala Leu Ala Asn> 580 590 600 610 620 TCA GTT AAA GAA CTT ACA AAT CCT GTT GTG GCA GAA AGT CCA AAA AAA AGT CAA TIT CTT GAA TGT TTA GGA CAA CAC CGT CTT TCA GGT TTT TTT Ser Val Lys Glu Leu Thr Asn Pro Val Val Ala Glu Ser Pro Lys Lys> 630 CCT TAA

FIGURE 13 (2 of 2)

WO 95/12676 PCT/US94/12352



24/133

OspC-PKO Sequence Range: 1 to 639

20 ATG AAA AAG AAT ACA TTA AGT GCG ATA TTA ATG ACT TTA TTT TTA TTT TAC TTT TTC TTA TGT AAT TCA CGC TAT AAT TAC TGA AAT AAA AAT AAA Met Lys Lys Asn Thr Leu Ser Ala Ile Leu Met Thr Leu Phe Leu Phe> 70 60 80 90 50 ATA TCT TGT AGT AAT TCA GGG AAA GGT GGG GAT TCT GCA TCT ACT AAT TAT AGA ACA TCA TTA AGT CCC TTT CCA CCC CTA AGA CGT AGA TGA TTA Ile Ser Cys Ser Asn Ser Gly Lys Gly Gly Asp Ser Ala Ser Thr Asn> 140 100 110 120 130 CCT GCT GAC GAG TCT GCG AAA GGG CCT AAT CTT ACA GAA ATA AGC AAA GGA CGA CTG CTC AGA CGC TTT CCC GGA TTA GAA TGT CTT TAT TCG TTT Pro Ala Asp Glu Ser Ala Lys Gly Pro Asn Leu Thr Glu-Ile Ser Lys> 160 170 180 AAA ATT ACA GAT TCT AAT GCA TTT GTA CTT GCT GTT AAA GAA GTT GAG TTT TAA TGT CTA AGA TTA CGT AAA CAT GAA CGA CAA TTT CTT CAA CTC Lys Ile Thr Asp Ser Asn Ala Phe Val Leu Ala Val Lys Glu Val Glu> 210 220 · 230 ACT TTG GTT TTA TCT ATA GAT GAA CTT GCT AAG AAA GCT ATT GGT CAA TGA AAC CAA AAT AGA TAT CTA CTT GAA CGA TTC TTT CGA TAA CCA GTT Thr Leu Val Leu Ser Ile Asp Glu Leu Ala Lys Lys Ala Ile Gly Gln> 260 270 280 AAA ATA GAC AAT AAT AAT GGT TTA GCT GCT TTA AAT AAT CAG AAT GGA TTT TAT CTG TTA TTA TTA CCA AAT CGA CGA AAT TTA TTA GTC TTA CCT Lys Ile Asp Asn Asn Asn Gly Leu Ala Ala Leu Asn Asn Gln Asn Gly> 290 300 310 320 330 TCG TTG TTA GCA GGA GCC TAT GCA ATA TCA ACC CTA ATA ACA GAA AAA AGC AAC AAT CGT CCT CGG ATA CGT TAT AGT TGG GAT TAT TGT CTT TTT Ser Leu Leu Ala Gly Ala Tyr Ala Ile Ser Thr Leu Ile Thr Glu Lys> 340 350 360 370 380 TTG AGT AAA TTG AAA AAT TTA GAA GAA TTA AAG ACA GAA ATT GCA AAG AAC TCA TTT AAC TTT TTA AAT CTT CTT AAT TTC TGT CTT TAA CGT TTC

Leu Ser Lys Leu Lys Asn Leu Glu Glu Leu Lys Thr Glu Ile Ala Lys>

OspC-PKO

420 430 410 390 GCT AAG AAA TGT TCC GAA GAA TTT ACT AAT AAA CTA AAA AGT GGT CAT CGA TTC TTT ACA AGG CTT CTT AAA TGA TTA TTT GAT TTT TCA CCA GTA Ala Lys Lys Cys Ser Glu Glu Phe Thr Asn Lys Leu Lys Ser Gly His> 470 480 450 440 GCA GAT CTT GGC AAA CAG GAT GCT ACC GAT GAT CAT GCA AAA GGA GCT CGT CTA GAA CCG TTT GTC CTA CGA TGG CTA CTA GTA CGT TTT CGT CGA Ala Asp Leu Gly Lys Gln Asp Ala Thr Asp Asp His Ala Lys Ala Ala> **5**20 510 500 490 ATT TTA AAA ACA CAT GCA ACT ACC GAT AAA GGT GCT AAA GAA TTT AAA TAA AAT TIT TGT GTA CGT TGA TGG CTA TIT CCA CGA TIT CIT AAA TIT Ile Leu Lys Thr His Ala Thr Thr Asp Lys Gly Ala Lys Glu Phe Lys> 570 560 550 540 530 GAT TTA TTT GAA TCA GTA GAA GGT TTG TTA AAA GCA GCT CAA GTA GCA CTA AAT AAA CTT AGT CAT CTT CCA AAC AAT TTT CGT CGA GTT CAT CGT Asp Leu Phe Glu Ser Val Glu Gly Leu Leu Lys Ala Ala Gln Val Ala> 620 600 590 580 CTA ACT AAT TCA GTT AAA GAA CTT ACA AGT CCT GTT GTA GCA GAA AGT . GAT TGA TTA AGT CAA TTT CTT GAA TGT TCA GGA CAA CAT CGT CTT TCA

Leu Thr Asn Ser Val Lys Glu Leu Thr Ser Pro Val Val Ala Glu Ser>

630

CCA AAA AAA CCT TAA GGT TTT TTT GGA ATT Pro Lys Lys Pro ***>

OspC-TRO

Sequence Range: 1 to 624

ATG AAA AAG AAT ACA TTA AGT GCG ATA TTA ATG ACT TTA TTT TTA TTT TAC TTT TTC TTA TGT AAT TCA CGC TAT AAT TAC TGA AAT AAA AAT AAA Het Lys Lys Asn Thr Leu Ser Ala Ile Leu Met Thr Leu Phe Leu Phe> 50 60 80 ATA TCT TGT AAT AAT TCA GGT GGG GAT TCT GCA TCT ACT AAT CCT GAT TAT AGA ACA TTA TTA AGT CCA CCC CTA AGA CGT AGA TGA TTA GGA CTA Ile Ser Cys Asn Asn Ser Gly Gly Asp Ser Ala Ser Thr Asn Pro Asp> 110 GAG TCT GCA AAA GGA CCT AAT CTT ACC GTA ATA AGC AAA AAA ATT ACA CTC AGA CGT TTT CCT GGA TTA GAA TGG CAT TAT TCG TTT TTT TAA TGT Glu Ser Ala Lys Gly Pro Asn Leu Thr Val Ile Ser Lys Lys Ile Thr> 160 180 GAT TCT AAT GCA TIT TTA CTG GCT GTG AAA GAA GTT GAG GCT TTG CTT CTA AGA TTA CGT AAA AAT GAC CGA CAC TTT CTT CAA CTC CGA AAC GAA Asp Ser Asn Ala Phe Leu Leu Ala Val Lys Glu Val Glu Ala Leu Leu> 200 210 220 230 TCA TCT ATA GAT GAA CTT TCT AAA GCT ATT GGT AAA AAA ATA AAA AAT AGT AGA TAT CTA CTT GAA AGA TTT CGA TAA CCA TTT TTT TAT TTT TTA Ser Ser Ile Asp Glu Leu Ser Lys Ala Ile Gly Lys Lys Ile Lys Asn> 250 260 270 280 GAT GGT ACT TTA GAT AAC GAA GCA AAT CGA AAC GAA TCA TTG ATA GCA CTA CCA TGA AAT CTA TTG CTT CGT TTA GCT TTG CTT AGT AAC TAT CGT Asp Gly Thr Leu Asp Asn Glu Ala Asn Arg Asn Glu Ser Leu Ile Ala> 290 300 310 320 GGA GCT TAT GAA ATA TCA AAA CTA ATA ACA CAA AAA TTA AGT GTA TTG CCT CGA ATA CTT TAT AGT TTT GAT TAT TGT GTT TTT AAT TCA CAT AAC Gly Ala Tyr Glu Ile Ser Lys Leu Ile Thr Gln Lys Leu Ser Val Leu> 350 360 370 AAT TCA GAA GAA TTA AAG AAA AAA ATT AAA GAG GCT AAG GAT TGT TCC TTA AGT CTT CTT AAT TTC TTT TTT TAA TTT CTC CGA TTC CTA ACA AGG Asn Ser Glu Glu Leu Lys Lys Lys Ile Lys Glu Ala Lys Asp Cys Ser>

OF E JCTOS HELDERS

OspC-TRO

420 430 410 400 390 GAA AAA TTT ACT ACT AAG CTA AAA GAT AGT CAT GCA GAG CTT GGT ATA CTT TTT AAA TGA TGA TTC GAT TTT CTA TCA GTA CGT CTC GAA CCA TAT Glu Lys Phe Thr Thr Lys Leu Lys Asp Ser His Ala Glu Leu Gly Ile> CAA AGC GTT CAG GAT GAT AAT GCA AAA AAA GCT ATT TTA AAA ACA CAT GTT TCG CAA GTC CTA CTA TTA CGT TTT TTT CGA TAA AAT TTT TGT GTA Gln Ser Val Gln Asp Asp Asn Ala Lys Lys Ala Ile Leu Lys Thr His> 520 490 GGA ACT AAA GAC AAG GGT GCT AAA GAA CTT GAA GAG TTA TTT AAA TCA CCT TGA TTT CTG TTC CCA CGA TTT CTT GAA CTT CTC AAT AAA TTT AGT Gly Thr Lys Asp Lys Gly Ala Lys Glu Leu Glu Glu Leu Phe Lys Ser> 550 560 570 540 530 CTA GAA AGC TTG TCA AAA GCA GCG CAA GCA GCA TTA ACT AAT TCA GTT GAT CTT TCG AAC AGT TTT CGT CGC GTT CGT CGT AAT TGA TTA AGT CAA Leu Glu Ser Leu Ser Lys Ala Ala Gln Ala Ala Leu Thr Asn Ser Val> 610 620 600 590 580 AAA GAG CTT ACA AAT CCT GTT GTG GCA GAA AGT CCA AAA AAA CCT TAA TIT CTC GAA TGT TTA GGA CAA CAC CGT CTT TCA GGT TTT TTT GGA ATT Lys Glu Leu Thr Asn Pro Val Val Ala Glu Ser Pro Lys Lys Pro ***>

P93

Sequence Range: 1 to 2102

30 10 ATG AAA AAA ATG TTA CTA ATC TTT AGT TTT TTT CTT ATT TTC TTG AAT TAC TIT TIT TAC AAT GAT TAG AAA TCA AAA AAA GAA TAA AAG AAC TTA Met Lys Lys Met Leu Leu Ile Phe Ser Phe Phe Leu Ile Phe Leu Asn> 90 .80 70 60 50 GGA TTT CCT GTT AGT GCA AGA GAA GTT GAT ,AGG GAA AAA TTA AAG GAC CCT AAA GGA CAA TCA CGT TCT CTT CAA CTA TGG-CTT TTT AAT TTC CTG Gly Phe Pro Val Ser Ala Arg Glu Val Asp Arg Glu Lys Leu Lys Asp> 140 130 120 110 100 TIT GIT AAT ATG GAT CIT GAG TIT GTA AAT TAT AAA GGC CCT TAT GAT AAA CAA TTA TAC CTA GAA CTC 1. A CAT TTA ATA TTT- CCG GGA ATA CTA Phe Val Asn Met Asp Leu Glu Phe Val Asn Tyr Lys Gly Pro Tyr Asp> 190 180 170 160 150 TCT ACA AAT ACA TAT GAA CAA ATA GTG GGT ATT GGG GAG TTT TTA GCA AGA TGT TTA TGT ATA CTT GTT TAT CAC CCA TAA CCC CTC AAA AAT CGT Ser Thr Asn Thr Tyr Glu Gln Ile Val Gly Ile Gly Glu Phe Leu Ala> 230 220 210 200 AGA CCG TTG ACC AAT TCC AAT AGC AAC TCA AGT TAT TAT GGT AAA TAT TCT GGC AAC TGG TTA AGG TTA TCG TTG AGT TCA ATA ATA CCA TTT ATA Arg Pro Leu Thr Asn Ser Asn Ser Asn Ser Ser Tyr Tyr Gly Lys Tyr> 280 270 260 TTT ATT AAT AGA TTT ATT GAT GAT CAA GAT AAA AAA GCA AGC GTT GAT AAA TAA TTA TCT AAA TAA CTA CTA GTT CTA TTT TTT CGT TCG CAA CTA Phe Ile Asn Arg Phe Ile Asp Asp Gln Asp Lys Lys Ala Ser Val Asp> 330 310 320 300 290 GTT TTT TCT ATT GGT AGT AAG TCA GAG CTT GAC AGT ATA TTG AAT TTA CAA AAA AGA TAA CCA TCA TTC AGT CTC GAA CTG TCA TAT AAC TTA AAT Val Phe Ser Ile Gly Ser Lys Ser Glu Leu Asp Ser Ile Leu Asn Leu> 370 360 350 340 AGA AGA ATT CTT ACA GGG TAT TTA ATA AAG TCT TTC GAT TAT GAC AGG

FIGURE 16 (1 of 5)

TCT TCT TAA GAA TGT CCC ATA AAT TAT TTC AGA AAG CTA ATA CTG TCC Arg Arg Ile Leu Thr Gly Tyr Leu Ile Lys Ser Phe Asp Tyr Asp Arg>

400 TCT AGT GCA GAA TTA ATT GCT AAG GTT ATT ACA ATA TAT AAT GCT GTT AGA TCA CGT CTT AAT TAA CGA TTC CAA TAA TGT TAT ATA TTA CGA CAA Ser Ser Ala Glu Leu Ile Ala Lys Val Ile Thr Ile Tyr Asn Ala Val> 450 460 470 TAT AGA GGA GAT TTG GAT TAT TAT AAA GGG TTT TAT ATT GAG GCT GCT ATA TCT CCT CTA AAC CTA ATA ATA TTT CCC AAA ATA TAA CTC CGA CGA Tyr Arg Gly Asp Leu Asp Tyr Tyr Lys Gly Phe Tyr Ile Glu Ala Ala> 500 490 520 TTA AAG TCT TTA AGT AAA GAA AAT GCA GGT CTT TCT AGG GTT TAT AGT AAT TTC AGA AAT TCA TTT CTT TTA CGT CCA GAA AGA TCC CAA ATA TCA Leu Lys Ser Leu Ser Lys Glu Asn Ala Gly Leu Ser Arg Val Tyr Ser> 530 540 550 560 CAG TGG GCT GGA AAG ACA CAA ATA TTT ATT CCT CTT AAA AAG GAT ATT GTC ACC CGA CCT TTC TGT GTT TAT AAA TAA GGA GAA TTT TTC CTA TAA Gln Trp Ala Gly Lys Thr Gln Ile Phe Ile Pro Leu Lys Lys Asp Ile> 580 590 600 610 TTG TCT GGA AAT ATT GAG TCT GAC ATT GAT ATT GAC AGT TTA GTT ACA AAC AGA CCT TTA TAA CTC AGA CTG TAA CTA TAA CTG TCA AAT CAA TGT Leu Ser Gly Asn Ile Glu Ser Asp Ile Asp Ile Asp Ser Leu Val Thr> 630 640 650 €€0 GAT AAG GTG GTG GCA GCT CTT TTA AGT GAA AAT GAA GCA GGT GTT AAC CTA TTC CAC CAC CGT CGA GAA AAT TCA CTT TTA CTT CGT CCA CAA TTG Asp Lys Val Val Ala Ala Leu Leu Ser Glu Asn Glu Ala Gly Val Asn> 690 700 TTT GCA AGA GAT ATT ACA GAT ATT CAA GGC GAA ACT CAT AAG GCA GAT AAA CGT TCT CTA TAA TGT CTA TAA GTT CCG CTT TGA GTA TTC CGT CTA Phe Ala Arg Asp Ile Thr Asp Ile Gln Gly Glu Thr His Lys Ala Asp> 740 750 760 CAA GAT AAA ATT GAT ATT GAA TTA GAC AAT ATT CAT GAA AGT GAT TCC GTT CTA TTT TAA CTA TAA CTT AAT CTG TTA TAA GTA CTT TCA CTA AGG Gln Asp Lys Ile Asp Ile Glu Leu Asp Asn Ile His Glu Ser Asp Ser> 770 790 800 810 AAT ATA ACA GAA ACT ATT GAA AAT TTA AGG GAT CAG CTT GAA AAA GCT TTA TAT TGT CTT TGA TAA CTT TTA AAT TCC CTA GTC GAA CTT TTT CGA Asn Ile Thr Glu Thr Ile Glu Asn Leu Arg Asp Gln Leu Glu Lys Ale>

FIGURE 16 (2 of 5)

820 830 840 850 860.

ACA GAT GAA GAG CAT AAA AAA GAG ATT GAA AGT CAG GTT GAT GCT AAA TGT CTA CTT CTC GTA TTT TTT CTC TAA CTT TCA GTC CAA CTA CGA TTT Thr Asp Glu Glu His Lys Lys Glu Ile Glu Ser Gln Val Asp Ala Lys>

870 880 890 900 910

AAG AAA CAA AAG GAA GAG CTA GAT AAA AAG GCA ATA AAT CTT GAT AAA TTC TTT GTT TTC CTT CTC GAT CTA TTT TTC CGT TAT TTA GAA CTA TTT Lys Lys Glu Lys Glu Glu Leu Asp Lys Lys Ala Ile Asn Leu Asp Lys

920 930 940 950 960

GCT CAG CAA AAA TTA GAT TCT GCT GAA GAT AAT TTA GAT GTT CAA AGA CGA GTC GTT TTT AAT CTA AGA CGA CTT CTA TTA AAT CTA CAA GTT TCT Ala Gln Gln Lys Leu Asp Ser Ala Glu Asp Asn Leu Asp Val Gln Arg>

970 980 990 1000

AAT ACT GTT AGA GAG AAA ATT CAA GAG GAT ATT AAC GAA ATT AAC AAG TTA TGA CAA TCT CTC TTT TAA GTT CTC CTA TAA TTG CTT TAA TTG TTC Asn Thr Val Arg Glu Lys Ile Gln Glu Asp Ile Asn Glu Ile Asn Lys>

1010 1020 1030 1040 1050

GAA AAG AAT TTA CCA AAG CCT GGT GAT GTA AGT TCT CCT AAA GTT GAT CTT TTC TTA AAT GGT TTC GGA CCA CTA CAT TCA AGA GGA TTT CAA CTA Glu Lys Asn Leu Pro Lys Pro Gly Asp Val Ser Ser Pro Lys Val Asp>

1060 1070 1080 1090 1100

AAG CAA CTA CAA ATA AAA GAG AGC CTG GAA GAT TTG CAG GAG CAG CTT TTC GTT GAT GTT TAT TTT CTC TCG GAC CTT CTA AAC GTC CTC GTC GAA Lys Gln Leu Gln Ile Lys Glu Ser Leu Glu Asp Leu Gln Glu Gln Leu>

1110 1120 1130 1140 1150

AAA GAA ACT GGT GAT GAA AAT CAG AAA AGA GAA ATT GAA AAG CAA ATT TTT CTT TGA CCA CTA CTT TTA GTC TTT TCT CTT TAA CTT TTC GTT TAA Lys Glu Thr Gly Asp Glu Asn Gln Lys Arg Glu Ile Glu Lys Gln Ile>

1160 1170 1180 1190 1200

GAA ATC AAA AAA AGT GAT GAA AAG CTT TTA AAA AGT AAA GAT GAT AAA CTT TAG TTT TTT TCA CTA CTT TTC GAA AAT TTT TCA TTT CTA CTA TTT Glu Ile Lys Lys Ser Asp Glu Lys Leu Leu Lys Ser Lys Asp Asp Lys>

1210 1220 1230 1240

GCA AGT AAA GAT GGT AAA GCC TTG GAT CTT GAT CGA GAA TTA AAT TCT CGT TCA TTT CTA CCA TTT CGG AAC CTA GAA CTA GCT CTT AAT TTA AGA Ala Ser Lys Asp Gly Lys Ala Leu Asp Leu Asp Arg Glu Leu Asn Ser>

FIGURE 16 (3 of 5)

1290 1260 1270 1260 1250 AAA GCT TCT AGC AAA GAA AAA AGT AAA GCC AAG GAA GAA GAA ATA ACC TTT CGA AGA TCG TTT CTT TTT TCA TTT CGG TTC CTT CTT TAT TGG Lys Ala Ser Ser Lys Glu Lys Ser Lys Ala Lys Glu Glu Glu Ile Thr> 1330 1320 1310 1300 AAG GGT AAG TCA CAG AAA AGC TTA GGC GAT TTG AAT AAT GAT GAA AAT TTC CCA TTC AGT GTC TTT TCG AAT CCG CTA AAC TTA TTA CTA CTT TTA Lys Gly Lys Ser Gln Lys Ser Leu Gly Asp Leu Asn Asn Asp Glu Asn> 1370 . `---1380 1360 1350 CTT ATG ATG CCA GAA GAT CAA AAA TTA CCT GAG GTT AAA AAA TTA GAT GAA TAC TAC GGT CTT CTA GTT TTT AAT GGA CTC CAA TTT TTT AAT CTA Leu Met Met Pro Glu Asp Gln Lys Leu Pro Glu Val Lys Lys Leu Asp> 1440 1430 1420 1410 1400 AGC AAA AAA GAA TTT AAA CCT GTT TCT GAG GTT GAG AAA TTA GAT AAG TCG TTT TTT CTT AAA TTT GGA CAA AGA CTC CAA CTC TTT AAT CTA TTC Ser Lys Lys Glu Phe Lys Pro Val Ser Glu Val Glu Lys Leu Asp Lys> 1480 1470 1460 1450 ATT TTC AAG TCT AAT AAC AAT GTT GGA GAA TTA TCA CCG TTA GAT AAA TAA AAG TTC AGA TTA TTG TTA CAA CCT CTT AAT AGT GGC AAT CTA TTT Ile Phe Lys Ser Asn Asn Asn Val Gly Glu Leu Ser Pro Leu Asp Lys> 1530 1520 1510 1500 1490 TCT TCT TAT AAA GAC ATT GAT TCA AAA GAG GAG ACA GTT AAT AAA GAT AGA AGA ATA TTT CTG TAA CTA AGT TTT CTC CTC TGT CAA TTA TTT CTA Ser Ser Tyr Lys Asp Ile Asp Ser Lys Glu Glu Thr Val Asn Lys Asp> 1570 1560 1550 1540 GTT AAT TTG CAA AAG ACT AAG CCT CAG GTT AAA GAC CAA GTT ACT TCT CAA TTA AAC GTT TTC TGA TTC GGA GTC CAA TTT CTG GTT CAA TGA AGA Val Asn Leu Gln Lys Thr Lys Pro Gln Val Lys Asp Gln Val Thr Ser> 1620 1610 1600 1590 TTG AAT GAA GAT TTG ACT ACT ATG TCT ATA GAT TCC AGT AGT CCT GTA AAC TTA CTT CTA AAC TGA TGA TAC AGA TAT CTA AGG TCA TCA GGA CAT . Leu Asn Glu Asp Leu Thr Thr Met Ser Ile Asp Ser Ser Ser Pro Val> 1670 1660 1650 1640 TIT TTA GAG GTT ATT GAT CCA ATT ACA AAT TTA GGA ACT CTT CAA CTT AAA AAT CTC CAA TAA CTA GGT TAA TGT TTA LAT CCT TGA GAA GTT GAA Phe Leu Glu Val Ile Asp Pro Ile Thr Asn Leu Gly Thr Leu Gln Leu>

1690 1700 1710 1720

ATT GAT TTA AAT ACT GGT GTT AGG CTT AAA GAA AGC ACT CAG CAA GGC TAA CTA AAT TTA TGA CCA CAA TCC GAA TTT CTT TCG TGA GTC GTT CCG Ile Asp Leu Asn Thr Gly Val Arg Leu Lys Glu Ser Thr Gln Gln Gly>

1730 1740 1750 1760 1770

ATT CAG CGG TAT GGA ATT TAT GAA CGT GAA AAA GAT TTG GTT GTT ATT TAA GTC GCC ATA CCT TAA ATA CTT GCA CTT TTT CTA AAC CAA CAA TAA Ile Gln Arg Tyr Gly Ile Tyr Glu Arg Glu Lys Asp Leu Val Val Ile>

1780 1790 1800 1810 1820

AAA ATG GAT TCA GGA AAA GCT AAG CTT CAG ATA CTT GAT AAA CTT GAA TTT TAC CTA AGT CCT TTT CGA TTC GAA GTC TAT GAA CTA TTF GAA CTT Lys Met Asp Ser Gly Lys Ala Lys Leu Gln Ile Leu Asp Lys Leu Glu>

1830 1840 1850 1860 1870

AAT TTA AAA GTG GTA TCA GAG TCT AAT TTT GAG ATT AAT AAA AAT TCA TTA AAT TTT CAC CAT AGT CTC AGA TTA AAA CTC TAA TTA TTT TTA AGT Asn Leu Lys Val Val Ser Glu Ser Asn Phe Glu Ile Asn Lys Asn Ser>

1880 1890 1900 1910 1920

TCT CTT TAT GTT GAT TCT AAA ATG ATT TTA GTA GCT GTT AGG GAT AAA AGA GAA ATA CAA CTA AGA TTT TAC TAA AAT CAT CGA CAA TCC CTA TTT Ser Leu Tyr Val Asp Ser Lys Met Ile Leu Val Ala Val Arg Asp Lys>

1930 1940 1950 1960

1970 1980 1990 2000 2010

GAT GAG TTT ATT CTT TCA GAG AAT AAA ATT ATG CCT TTT ACT AGC TTT CTA CTC AAA TAA GAA AGT CTC TTA TTT TAA TAC GGA AAA TGA TCG AAA ASP Glu Phe Ile Leu Ser Glu Asn Lys Ile Met Pro Phe Thr Ser Phe>

2020 2030 2040 2050 2060

TCT GTG AGA AAA AAT TTT ATT TAT TTG CAA GAT GAG TTT AAA AGT CTA AGA CAC TCT TTT TTA AAA TAA ATA AAC GTT CTA CTC AAA TTT TCA GAT Ser Val Arg Lys Asn Phe Ile Tyr Leu Gln Asp Glu Phe Lys Ser Leu>

2070 2080 2090 2100

GTT ATT TTA GAT GTA AAT ACT TTA AAA AAA GTT AAG TA CAA TAA AAT CTA CAT TTA TGA AAT TTT TTT CAA TTC AT Val lle Leu Asp Val Asn Thr Leu Lys Lys Val Lys Xxx>

FIGURE 16 (5 of 5)

p93 - K48

			4			
1	ATGAAAAAAT	TGTTACTAAT	CTTTAGTTTT	TTTCTTATTT	CTTTGAATGG	ATTICCTCTT
63	P PARKEY PCCC	AACTTCATAA	GGAAAAATTA	AAGGATTTTG	TTAATATGGA	TCITGAGTTT
121	ATATA A ATA	A A COTY COTTA	TGATTCTACA	AATACATATG	AACAAATAGT	AGGIATIGGT
101	CyCatalalalaby C	CAAGACCATT	GATTAATTCC	AATAGCAACT	CAATITATTA	TGGTAAATAT
241	ፈጥፈ ልጥጥ ልጥጥ	CATTTATTGA	TGATCAAGAT	AAAAAAGCAA	GCGTTGATGT	TTTTTCTATT
201	CCTACTACCT	CACAGCTTGA	CAGTATATTG	AATCTAAGAA	GAATTCITAC	AGGGTATTTG
261	ALL DE VILLE	TYCATTATGA	AAGATCTAGT	GCTGAATTAA	TTGCTAAGGT	TATTACAATA
421	CATABITECTE	TTTATAGAGG	GGATTTAAAT	TATTATAAAG	AGGITTATAT	TGAGGCTGCT
701	JALLAND & VALLE	TAACTAAAGA	AAATGCAGGT	CTTTCTAGAG	TGTACAGTCA	ATGGGCTGGA
CAT	AAAAAAAA	TATTTATTCC	TCTTAAAAAG	TATTTTTAT	CIGGAAAAGT	TGAGTCTGAC
601	STPP APPAPPA	ACAGTTTGGT	TACAGATAAG	GTTGTGGCAG	-CICITITAAG	CGAGAATGAA
661	GCAGGTGTTA	ACTITICAAG	AGATATTACA	GATATTCAAG	GCGAAACTCA	TAAAGCAGAT
221	CAAGATAAAA	TYCATATICA	ATTAGATAAT	GTTCATAAAA	GTGATTCCAA	TATAACAGAG
701	ACTATION CA	ATTTAAGAGA	TCAGCTTGAA	AAGGCTACAG	ATGAAGAGCA	TAGAAAAGAG
0/1	ATTGAAAGTC	ACCTTGATGC	TAAAAAGAAA	CAAAAAGAAG	AACTAGATAA	AAAGGCAATC
001	ATTACOUTY AT A	AAGCCCAACA	AAAATTAGAT	TCTTCTGAAG	ATAATTTAGA	TATTCAAAGG
961	CATACTCTTA	GAGAGAAGAT	TCAAGAGGAT	ATTGACGAGA	TTAATAAAGA	AAAGAATTTG
1021	CCDDDACCTG	GTGATGTAAG	TTCTCCTAAA	GTTGATAAGC	AGCTACAAAT	AAAAGAGAGT
1081	CTACAACACT	TGCAGGAACA	GCTTAAAGAA	ACTAGCGATG	AAAATCAAAA	AAGAGAAATT
11/1	GAAAAGCAAA	TTGAAATCAA	AAAAAGTGAT	GAAGAACTTT	TAAAAAGTAA	AGATCCTAAA
1201	GCATTAGATC	TTAATGGAGA	TTTAAATTCT	AAAGTTTCTA	GTAAAGAAAA	AATTAAAGGC
1261	AAAGAAGGAG	AAATAGTCAA	AGAGGAATCA	AAGGCAAGTT	'TAGCTGATTT	GAATAATGAC
1321	CANANTYTTA	TGAGGCCGGA	AGATCAAAAA	TTATCTGAGG	TTAAAAATK	AGATAGTAAA
1381	AATTTAA	AACCTGTTTC	TGAGATTGAG	AGAGTAAATG	AAATTTCGAA	GTCTAACAAC
1441	AATCACATTA	GTGAATCATC	ACCATTATAT	· AAGCCTTCTT	' ATAGCGATAT	GGATTCAAAA
1501	CACCGTATAG	ATAATAAAGA	TGTTAACTTG	CAAGAAACCA	. AGTCTCAAAC	TAAAAGTCAA
1561	CCTACTTCTT	TAAATCAAGA	TTTGACTACT	' ATGTCTATAG	ATTCTAGTAA	TCCTCTATTT
1621	TTAGAGGTTA	TIGATCCTAT	TACAAATTTA	GGAACGCTTC	: AACTTATTGA	TTTGAATACC
1681	CCTCTTAGAC	TTAAAGAAAC	TACTCAGCAA	GGCATTCAGC	GGTATGGAAT	TTATGAACGT
1741	CAAAAAGATT	TAGTIGITAT	TAAAATGGAI	' TCAGGAAAAC	CCAAGCTTCA	AATACTTAAT
1801	AAACTTGAGA	ATTTAAAAGT	CATATCGGAC	TCTAATTITC	agattaataa	ANATTCATCT
1861	THEST ATTENT	ACTCTAAAA?	GATTTTAGT!	A GTTGTGAGAC	ATAGTGGTAA	TGTTTGGAGA
1921	TTCCCTAAAT	TTTCTCCTA	CAAATTTAAA A	CAGTTTATTY 1	: TTTCAGAGAA	TAAAATTTTG
1981	CCTTTTACTA	GCTTTTCTG	r gagaaagaa?	TATTATTT 1	r igcaggatga	GTTTAA&AGT
204	CTTATTACT	TAGATGTAA	A TACTITAAA	AAAGTTAAG	A 7	

p93 - BO

	ATGAAAAAA					
	AATGCAAGGG					
121	GTTAATTACA	AGGGTCCTTA	TGATTCTACA	GATACATATG	AACAAATAGT	AGGTATTGGG
181	GAGTTTTTAG	CAAGGCCGTT	GAACAATTCC	AATAGTAATT	CAAGTTATTA	TGGTAAATAT
241	TTTGTTAATA	GATTTATTGA	CGATCAAGAT	AAAAAAGCAA	GTGTTGATAT	TTTTTCTATT
301	GGTAGTAAGT	CAGAGCTTGA	TAGTATATTA	AATCTAAGAA	GAATTCTTAC	AGGGTATTTA
361	ATGAAGTCTT	TTGATTATGA	GAGGTCTAGT	GCGGAATTAA	TTGCTAAAGC	TATTACAATA
421	TATAATGCTG	TTTATAGAGG	AGATTTAGAT	TATTACAAAG	AGTTTTATAT	TGAGGCTTCT
481	TTGAAGTCTT	TGACTAAAGA	AAATGCAGGT	CTTTCTAGGG	TGTACAGTCA	ATGGGCTGGG
541	AAGACACAAA	TATTTATTCC	TCTTAAAAAG	TATTTTATAA	CTGGAAATGT	TGAGTCTGAC
601	ATTGATATTG	ATAGTTTGGT	TACAGATAAG	GTGGTGGCAG	CTCTTTTAAG	TGAGAATGAA
661	TCAGGTGTTA	ACTITICAAG	AGATATTACA	GACATTCAAG	GCGAÁACTCA	TAAAGCAGAT
	CAAGATAAAA					
781	ACTATTGAGA	ATTTAAGGGA	TCAGCTTGAA	AAAGCTACAG	ATGAAGAGCA	TAAAAAAGAG
841	ATTGAAAGTC	AGGTTGATGC	TAAAAAGAAA	CAAAAGGAAG	AATTAGATAA	AAAGGCAATT
901	GATCTTGATA	AAGCTCAACA	AAAATTAGAT	TTTGCTGAAG	ATAATCTAGA	TATTCAAAGG
	GATACTGTTA					
	CCAAAGCCTG					
1081	CTAGAAGATT	TGCAAGAGCA	GCTTAAAGAA	GCTAGTGATG	AAAATCAAAA	AAGAGAAATA
	GAAAAGCAAA					
	GCATTAGATC					
	GAAGAAGAGG					
	AAAGTAGATA					
	GAGCCTTCTT					
	CAAAAAACTA					
	GTGTCTATAG					
	GGAACGCTTC					
	GGTATTCAGC					
	TCAGGAAAAG					
	TCTAATTTTG					
	GTTGTTAAGG					
	GAATTTATTC					
	TTATTTATT		ACTTAAAAGC	TTAGTTACTT	TAGATGTAAA	TACTTTAAAA
1981	AAAGTTAAGT	A				

p93 - p1RO

1	ATGAAAAAA	delicate of par	Cutatate Valadata	datab datab datab	CALALACY PARCO	A TVDDCCCccccc
	AATGCAAGGG					
	GTAAACTATA					
	GAGTTTTTAG					
	TTTATTAATA					
241	AGTAGTAAGT	CALITATION	CONTONIONI	AAAAAAAAAAAAA	COLIGATO!	TITTICTATT
301	ATAAAGTCTT	CACAGCI IOA	A N C N TO CT N C TO CT N	WILLIAM CAM	GAATICITAC	AGGGTATTTG
701	CATAATGCTG	TIGATIATOA	WAGAICIAGI	GCIGAATTAA	TIGCCAAGGI	TATTACAATA
401	TTAAAGTCTT	TAACTAAAGA	WWW.IOCWOOT.	CITICIAGAG	TGTACAGTCA	ATGGGCTGGA
241	AAGACACAAA	TATTTATTCC	TCTTAMAMAG	AATATTTAT	CIGGAAAAAT	TGAGTCTGAC
	ATTGATATTG					
	GCAGGTGTTA					
721	CAAGATAAAA	TIGATATIGA	ATTAGATAAT	GITCATGAAA	GTGATTCCAA	TATAACAGAA
781	ACTATTGAGA	ATITAAGAGA	TCAGCTTGAA	AAGGCTACAG	ATGAAGAGCA	TAGAAAAGAG
841	ATTGAAAGTC	AAGTTGATGC	TAAAAAGAAA	CAAAAAGAAG	AACTAGATAA	AAAGGCAATC
	GATCTTGATA					
	GATACTGTTA					
	CCAAAACCTG					
1081	CTAGAAGACT	TGCAGGAGCA	GCTTAAAGAA	ACTAGCGATG	AAAATCAAAA	AAGAGAAATT
1141	GAAAAGCAAA	TTGAAATCAA	AAAAAGTGAT	GAAGAACTTT	TAAAAAGCAA	AGATCCTAAA
	GCATTAGATC					
	AAAGAAAAAG					
	GAAACCCTTA					
1381	AATTTAA	AACCTGTTTC	TGAGATTGAG	AGAGTAAATG	AAATTTCAAA	GTCTAACAAC
	AATGAGGTTA					
1501	GAGGTTGTAG	ATAATAAAGA	TGTTAATTTG	CAAGAAACCA	AGCCTCAAGC	TAAAAGTCAA
1561	TCTACTTCTT	TAAATCAAGA	TTTGATTACT	ATGTCTATAG	ATTCTAGTAA	TCCTGTATTT
1621	TTAGAGGTTA	TTGATCCTAT	TACAAATTTA	GGAATGCTTC	AACTTATTGA	TTTAAATACT
1681	GGTGTTAGAC	TTAAAGAAAG	CACTCAGCAA	GGCATTCAGC	GTTATGGAAT	TTATGAACGT
1741	GAAAAAGATT	TAGTTGTTAT	TAAAATGGAT	TCAGGAAAAG	CTAAGCTTCA	AATACTTAAT
1801	AAACTTGAGA	ATTTAAAAGT	GATATCAGAG	TCTAATTTTG	AGATTAATAA	AAATTCATCT
1861	CTTTATGTTG	ACTCTAAAAT	GATTTTAGTA	GCTGTGAAAG	ATAGTGGTAA	TGTTTGGAGA
1921	TTGGCTAAAT	TTTCTCCTAA	AAATTTAGAT	GAGTTTATTC	TTTCAGAGAA	TAAAATTTTG
	CCTTTTACTA					GTTTAAAAGT
2041	CTTATTACTT	TAGATGTAAA	TACTTTAAAA	AAAGTTAAGT	A	

p93 - pGau

	_			•		
1	ATGAAAAAA	TGTTACTAAT	CTTTAGTTTT	TTTCTTGTTT	TTTTAAATGG	ATTTCCTCTT
61	AATGCAAGGG	AAGTTGATAA	GGAAAAATTA	AAGGACTTTG	TTAATATGGA	TCTTGAATTT
		AGGGTCCTTA				
181	GAGTTTTTAG	CAAGGCCGTT	GATCAATTCC	AATAGTAATT	CAAGTTATTA	TGGTAAATAT
241	TTTGTTAATA	GATTTATTGA	CGATCAAGAT	AAAAAAGCAA	GTGTTGATAT	TTTTTCTATT
301	GGTAGTAAGT	CAGAGCTTGA	TAGTATATTA	AATCTAAGAA	GAATTCTTAC	AGGGTATTTA
361	ATGAAGTCTT	TTGATTATGA	GAGGTCTAGT	GCGGAATTAA	TTGCTAAAGC	TATTACAATA
		TTTATAGAGG				
481	TTGAAGTCTT	TGACTAAAGA	AAATGCAGGT	CTTTCTAGGG	TGTACAGTCA	ATGGGCTGGG
541	AAGACACAAA	TATTTATTCC	TCTTAAAAAG	TATTTTTAT	CTGGAAATGT	TGAGTCTGAC
601	ATTGATATTG	ATAGTTTGGT	TACAGATAAG	GTGGTGGCAG	CTCTTTTAAG	TGAGAATGAA
661	TCAGGTGTTA	ACTITICCAAG	AGATATTACA	GACATTCAAG	GCGAAACTCA	TAAAGCAGAT
721	CAAGATAAAA	TTGATATTGA	ATTAGATAAT	ATTCATGAAA	GTGATTCCAA	TATAACAGAA
781	ACTATTGAGA	ATTTAAGGGA	TCAGCTTGAA	AAAGCTACAG	ATGAAGAGCA	TAAAAAAGAG
841	ATTGAAAGTC	AGGTTGATGC	TAAAAAGAAA	CAAAAGGAAG	AATTAGATAA	AAAGGCAATT
		AAGCTCAACA				
		GAGAGAAGCT				
		GTGATGTAAG				
1081	CTGGAAGATT	TGCAGGAGCA	GCTTAAAGAA	ACTGGTGATG	AAAATCAGAA	AAGAGAAATT
1141	GAAAAGCAAA	TTGAAATCAA	AAAAAGTGAT	GAAAAGCTTT	TAAAAAGTAA	AGATGATAAA
1201	GCAAGTAAAG	ATGGTAAAGC	CTTGGATCTT	GATCGAGAAT	TAAATTCTAA	AGCTTCTAGC
1261	AAAGAAAAA	GTAAAGCCAA	GGAAGAAGAA	ATAACCAAGG	GTAAGTCACA	GAAAAGCTTA
1321	GGCGATTTGA	ATAATGATGA	AAATCTTATG	ATGCCAGAAG	ATCAAAAATT	ACCTGAGGTT
1381	AAAAAATTAG	ATAGCAAAAA	AGAATTTAAA	CCTGTTTCTG	AGGTTGAGAA	ATTAGATAAG
1441	ATTITCAAGT	CTAATAACAA	TGTTGGAGAA	TTATCACCGT	TAGATAAATC	TTCTTATAAA
1501	GACATTGATT	CAAAAGAGGA	GACAGTTAAT	AAAGATGTTA	ATTTGCAAAA	GACTAAGCCT
1561	CAGGTTAAAG	ACCAAGTTAC	TTCTTTGAAT	GAAGATTTGA	CTACTATGTC	TATAGATTCC
1621	AGTAGTCCTG	TATTTTTAGA	GGTTATTGAT	CCAATTACAA	ATTTAGGAAC	TCTTCAACTT
1681	ATTGATTTAA	ATACTGGTGT	TAGGCTTAAA	GAAAGCACTC	AGCAAGGCAT	TCAGCGGTAT
1741	GGAATTTATG	AACGTGAAAA	AGATTIGGTT	GTTATTAAAA	TGGATTCAGG	AAAAGCTAAG
1801	CTTCAGATAC	TTGATAAACT	TGAAAATTTA	AAAGTGGTAT	CAGAGTCTAA	TTTTGAGATT -
1861	TTAAAAATA	CATCTCTTTA	TGTTGATTCT	AAAATGATTT	TAGTAGCTGT	TAGGGATAAA
1921	GATAGTAGTA	ATGATTGGAG	ATTGGCCAAA	TTTTCTCCTA	AAAATTTAGA	TGAGTTTATT
1981	CTTTCAGAGA	TATAAAATA	GCCTTTTACT	AGCTTTTCTG	TGAGAAAAA	TATTTATTTT
2041	TTGCAAGATG	AGTTTAAAAG	TCTAGTTATT	TTAGATGTAA	ATACTTTAAA	AAAAGTTAAG
2101	TAAAGCC					

 $\langle \cdot \rangle$

37/133

p93 - pKO

				Jalak Make	TTTTAAATGG .	ATTTCCTCTT
					TTTTAAATGG .	
61	AATGCAAGGG	AAGTTGATAA	GGAAAAATTA	AAGGACIIIG	TTAATATGGA AACAAATAGT	AGGTATTGGG
100	T GUALLY TALES	T TGCAAGATG	A ACTTAAAAG	C TTAGTTACT	T TAGATGTAAA	TACTTTAAAA
100	1 AAAGTTAAG	T A				
מכיו	1 WWWG111WG					

p93 - 25015

1	ATGAAAAAA	TGTTACTAAT	CTTTAGTTTT	TTTCTTATTT	TTTTGAATGG	ATTTCCTCTT
61	AATGCAAGGA	AAGTTGATAA	GGAAAAATTA	AAGGATTTTG	TTAATATGGA	TCTTGAGTTT
121	GTAAATTATA	AAGGTCCTTA	TGATTCTACA	AATACGTATG	AACAAATAGT	GGGTATTGGG
181	GAGTTTTTAG	CAAGACCGCT	GACCAATTCC	AATAGCAACT	CAAGTTATTA	TGGCAAATAT
241	TTTATTAATA	GATTTATTGA	TGATCAAGAT	AAAAAAGCAA	GTGTTGATGT	TTTTTCTATA
301	AGCAGCAAAT	CAGAGCTTGA	CAGTATATTG	AATTTAAGAA	GAATTCTTAC	AGGGTATATA
361	ATAAAGTCTT	TCGATTATGA	CAGGTCTAGT	GCAGAATTAA	TIGCTAAGGT	TATTACAATA
421	TATAATGCTG	TTTATAGAGG	AGATTTGGAT	TATTATAAAG	GGTTTTATAT	TGAGCCTGCT
		TAACTAAAGA				
541	AAGACTCAAA	TATTTATTCC	TCTTAAAAAG	GATATTTTGT	CTGGAAATAT	TGAATCTGAC
		ACAGTTTGGT				
		ACTITICAAG				
		TTGATACTGA				
		ATTTAAGGGA				
		AGGTTGATGC				
		AAGCTCAGCA				
		GAGAGAAAAT				
		GTGATGTAAG				
		TGCAGGAGCA				
		TTGAAATCAA				
		ATTATGAAGC				
		AGGTCAAGGA				
		ATGATAAAA				
		GTAAATTAGA				
		TTTCCAAGTC				
		ATGATATTGA				
		CTAAAGTTAA				
		TGTCTATAGA				
		GAACCCTGCA				
		GCATTCAGCG				
		CAGGAAAGGC				
		CTAATITTGA				
		CTGTTAGAGA				
		TGGATGAGTT		-		
		TATTTTAAAA		GATGAGCTTA	AAAATCTAGT	TATTTTAGAT
2101	GTAAATACTT	TAAAAAAAGT	TAAGTA			

K48 OSP A/PGAU OSP A FUSION

39/133

		10			2	0			30			40)		
	•		•	•		•		•	•		•		•	•	
ATG .															
TAC															
Met	Lys	Ľys	Tyr	Leu	Leu	Gly	Ile	Gly	Leu	Ile	Leu	Ala	Leu	Ile	λla>
50			60			7	0			80			90		
•		•	•		•		•	•	•	•,		•	•		•
TGT .	AAG	CAA	TAA	GTT	AGC	AGC	CTT	GAT	GAÀ	አጸጸ -	TAA	AGC	GTT	TCA	GTA.
ACA '	TTC	GTT	ATT	CAA	TCG	TCG	GAA	CTA	CTT	TTT	TTA	TCG	CAA	AGπ	CAT
Cys	Lys	Gln	Asn	Val	Ser	Ser	Leu	Asp	Glu	Lys	Asn	Ser	Val	Ser	Val>
10	O	•	1	10			120			13	0		1	140	
	*	•	_	•		•	•		•		•	•	-	•	
GAT	TTA	CCT	GGT	GGA	ATG	ACA	GTT	CTT	GTA	AGT	AAA	GAA	AAA	GAC	አሕአ
CTA	TAA	GGA	CCA	CCT	TAC	TGT	CAA	GAA	CAT	TCA	TTT	CTT	TTT	CTG	TTT
Asp	Leu	Pro	Gly	Gly	Met	Thr	Val	Leu	Val	Ser	Lys	Glu	Lys	Asp	Lys>
	150			16	50		1	70			180			19	? ∩
•	•		•	- `	•	•	-	•		•	•		•		•
GAC	GGT	AAA	TAC	AGT	CTA	GAG	GCA	ACA	GTA	GAC	AAG	CTT	GAG	CTT	AAA
														GAA	
															Lys>
	•	200			210			2:	20		:	230			240
•	•	•	•	•	•		•		•	•		•		•	•
GGA	ACT	TCT	GAT	AAA	AAC	AAC	GGT	TCT	GGA	ACA	CTT	GAA	GGT	GAA	AAA
CCT	TGA	AGA	CTA	TTT	TTG	TTG	CCA	AGA	CCT	TGT	GAA	CTT	CCA	CTT	TII
Gly	Thr	Ser	Asp	Lys	Asn	Asn	Gly	Ser	Gly	Thr	Leu	Glu	Gly	Glu	Lys>
		2	50		•	260			270			28	20		
	•	-													
ACT			•	•		•	•	•	•		•		•	•	
	GAC	AAA	AGT	AAA	GTA	AAA	TTA	• ACA	•	GCT	• GAT	•	•	AGT	CAA
TGA									ATT			GAC	• CTA	agt TCA	
	CTG	TTT	TCA	TTT	CAT	TTT	AAT	TGT	ATT TAA	CGA	CTA	GAC CTG	CTA GAT	TCA	
Thr	CTG	TTT	TCA Ser	TTT	CAT	TTT Lys	AAT Leu	TGT	ATT TAA Ile	CGA Ala	CTA	GAC CTG	CTA GAT Leu	TCA	GTT
	CTG	TTT	TCA	TTT	CAT	TTT Lys	AAT	TGT	ATT TAA Ile	CGA	CTA	GAC CTG	CTA GAT	TCA	GTT
Thr 290	CTG Asp	TTT Lys	TCA Ser	TTT Lys	CAT Val	TTT Lys	Leu 10	TGT Thr	ATT TAA Ile	CGA Ala 320	CTA Asp	GAC CTG Asp	CTA GAT Leu 330	TCA	GIT Glm>
Thr 290 ACT	CTG Asp	TTT Lys	TCA Ser 300 GAA	TTT Lys	CAT Val	TTT Lys 3	Leu 10 GAA	TGT Thr	ATT TAA Ile	CGA Ala 320 AAA	CTA Asp ACA	GAC CTG Asp	CTA GAT Leu 330 GTA	TCA Ser TCA	GIT Glm>
Thr 290 ACT TGA	CTG Asp AAA TTT	TTT Lys	TCA Ser 300 GAA CTT	TTT Lys ATT TAA	CAT Val TTC AAG	TTT Lys 3 AAA TTT	Leu 10 GAA CTT	TGT Thr GAT CTA	ATT TAA Ile GCC CGG	CGA Ala 320 AAA TTT	CTA Asp ACA TGT	GAC CTG Asp TTA	CTA GAT Leu 330 GTA CAT	TCA Ser TCA AGT	GTT Gln>
Thr 290 ACT TGA Thr	AAA TTT Lys	TTT Lys	TCA Ser 300 GAA CTT	TTT Lys ATT TAA Ile	CAT Val TTC AAG	TTT Lys 3 AAA TTT	AAT Leu 10 GAA CTT Glu	TGT Thr GAT CTA Asp	ATT TAA Ile GCC CGG	CGA Ala 320 AAA TTT Lys	ACA TGT	GAC CTG Asp TTA	CTA GAT Leu 330 GTA CAT Val	TCA Ser TCA AGT Ser	GTT Gln> AAA TTT
Thr 290 ACT TGA Thr	CTG Asp AAA TTT	TTT Lys	TCA Ser 300 GAA CTT	TTT Lys ATT TAA	CAT Val TTC AAG	TTT Lys 3 AAA TTT	Leu 10 GAA CTT	TGT Thr GAT CTA Asp	ATT TAA Ile GCC CGG	CGA Ala 320 AAA TTT Lys	CTA Asp ACA TGT	GAC CTG Asp TTA	CTA GAT Leu 330 GTA CAT Val	TCA Ser TCA AGT	GTT Gln> AAA TTT
Thr 290 ACT TGA Thr	AAA TTT Lys	TTT Lys TTT AAA Phe	TCA Ser 300 GAA CTT Glu	TTT Lys ATT TAA Ile 350	CAT Val TTC AAG Phe	TTT Lys 3 AAA TTT Lys	AAT Leu 10 GAA CTT Glu 360	TGT Thr GAT CTA Asp	ATT TAA Ile GCC CGG Ala	CGA Ala 320 AAA TTT Lys	ACA TGT Thr	GAC CTG Asp TTA AAT Leu	CTA GAT Leu 330 GTA CAT Val	TCA Ser TCA AGT Ser 380	GTT Glr> AAA TTT Lys>
Thr 290 ACT TGA Thr 3	AAA TTT Lys	TTT Lys TTT AAA Phe	TCA Ser 300 GAA CTT Glu	ATT TAA Ile	CAT Val TTC AAG Phe	TTT Lys AAA TTT Lys AAG	AAT Leu 10 GAA CTT Glu 360	TGT Thr GAT CTA Asp	ATT TAA Ile GCC CGG Ala	CGA Ala 320 AAA TTT Lys 3	ACA TGT Thr	GAC CTG Asp TTA AAT Leu	CTA GAT Leu 330 GTA CAT Val	TCA Ser TCA AGT Ser 380	GTT Gln> AAA TTT

FIGURE 23 (1 of 3)

K48 OSP A/ PGAU OSPA FUSION

400 430 390 AAG GGT GAA ACA TCT GAA AAA ACA ATA GTA AGA GCA AAT GGA ACC AGA TTC CCA CTT TGT AGA CTT TTT TGT TAT CAT TCT CGT TTA CCT TGG TCT Lys Gly Glu Thr Ser Glu Lys Thr Ile Val Arg Ala Asn Gly Thr Arg> 460 470 440 450 480 CTT GAA TAC ACA GAC ATA AAA AGC GAT GGA ,TCC GGA AAA GCT AAA GAA GAA CTT ATG TGT CTG TAT TTT TCG CTA CCT AGG CCT TTT CGA TTT CTT . Leu Glu Tyr Thr Asp Ile Lys Ser Asp Gly Ser Gly Lys Ala Lys Glu> 500 510 520 490 GTT TTA AAA GAC TTT ACT CTT GAA GGA ACT CTA GCT GCT GAC GGC AAA CAA AAT TIT CTG AAA TGA GAA CTT CCT TGA GAT CGA CGA CTG CCG TIT Val Leu Lys Asp Phe Thr Leu Glu Gly Thr Leu Ala Ala Asp Gly Lys> 550 560 570 530 540 ACA ACA TTG AAA GTT ACA GAA GGC ACT GTT GTT TTA AGC AAG AAC ATT TGT TGT AAC TTT CAA TGT CTT CCG TGA CAA CAA AAT TCG TTC TTG TAA Thr Thr Leu Lys Val Thr Glu Gly Thr Val Val Leu Ser Lys Asn Ile> 600 610 620 580 590 TTA AAA TCC GGA GAA ATA ACA GTT GCA CTT GAT GAC TCT GAC ACT ACT AAT TIT AGG CCT CIT TAT TGT CAA CGT GAA CTA CTG AGA CTG TGA TGA Leu Lys Ser Gly Glu Ile Thr Val Ala Leu Asp Asp Ser Asp Thr Thr> 650 660 630 640 CAG GCT ACT AAA AAA ACT GGA AAA TGG GAT TCA AAA ACT TCT ACT TTA' GTC CGA TGA TTT TTT TGA CCT TTT ACC CTA AGT TTT TGA AGA TGA AAT Gln Ala Thr Lys Lys Thr Gly Lys Trp Asp Ser Lys Thr Ser Thr Leu> 700 710 690 680 ACA ATT AGT GTT AAC AGC AAA AAA ACT. ACA CAA CTT GTG TTT ACT AAA TGT TAA TCA CAA TTG TCG TTT TTT TGA TGT GTT GAA CAC AAA TGA TTT Thr Ile Ser Val Asn Ser Lys Lys Thr Thr Gln Leu Val Phe Thr Lys> 750 760 740 730 CAA TAC ACA ATA ACT GTA AAA CAA TAC GAC TCC GCA GGT ACC AAT TTA GTT ATG TGT TAT TGA CAT TTT GTT ATG CTG AGG CGT CCA TGG TTA AAT Gln Tyr Thr Ile Thr Val Lys Gln Tyr Asp Ser Ala Gly Thr Asn Leu>

FIGURE 23 (2 of 3)

K48 OSPA / PGAU OSP A FUSION

770 780 790 800 810

GAA GGC ACA GCA GTC GAA ATT AAA ACA CTT GAT GAA CTT AAA AAC GCT CTT CCG TGT CGT CAG CTT TAA TTT TGT GAA CTA CTT GAA TTT TTG CGA Glu Gly Thr AIa Val Glu Ile Lys Thr Leu Asp Glu Leu Lys Asr. Ala>

820

TTA AAA TAA AAT TTT ATT Leu Lys ***>

FIGURE 23 (3 of 3)

B-31 OSP A /PGAU OSP A FUSION

	10		20		_	30			4	0		
ATG AAA	TAT AAA	TTA TTG	GGA A	ATA (GGT	CTA	ATA	TTA	GCC	TTA	ATA	GCA
TAC TTT	ATA TTT	AAT AAC	CCT :	TAT	CCA	GAT	TAT	TAA	CGG	TAA	TAT	CGT
Met Lys	Lys Tyr	Leu Leu	Gly :	Ile	Gly	Leu	Ile	Leu	Ala	Leu	Ile	Ala>
50	60		7	0			80			90		
•	• •	•	•	•	•		•		•	•		•
TGC AAG	CAA AAT	GTT AGC	AGC (CTT	GAT	GAA	WW.	AYC.	AGC	GCT	TCA	GTA
ACG TTC	GTT TTA	CAA TCG Val Ser	Ser 1	GAA '	yen Civ	Glu	Lvs	Asn	Ser	Ala	Ser	Val
Cys Lys	GIN ASH	Agt Set	Jer .	.	nop		-, -				٠.	4417
100	:	110	:	120		_	13	0	_	1	40	
	•	GAG ATG	*	~~~ ~~~~	ر محت	CTA	ACT	*	CAA	5.5.5	CNC.	
GAT TIG	CCT GGT	CTC TAC	TTT	CAA	GAA	CAT	TCA	TTT	CTT	TTT	CTG	TTT
Asp Leu	Pro Gly	Glu Met	Lys	Val	Leu	Val	Ser	Lys	Glu	Lys	Asp	Lys>
		1.60		,	70			180				
150	•	160	*	1			•	*		•	13	90 →
GAC GGT	AAG TAC	AGT CTA	AAG	GCA	ACA	GTA	GAC	AAG	ATT	GAG	CTA	AAA
CTG CCA	TTC ATG	TCA GAT	TTC	CGT	TGT	CAT	CTG	TTC	TAA	CTC	GAT	TTT
Asp Gly	Lys Tyr	Ser Leu	Lys	Ala	Thr	Val	Asp	Lys	Ile	Glu	Leu	Lys>
	200	210			22	20			230			240
•	•	• •		•		•	•		•		•	•
GGA ACT	TCT GAT	AAA GAC	TAA	GGT	TCT	GGA	GTG	CTT	GAA	GGT	ACA	AAA
CCT TGA	AGA CTA	TTT CTG	ATT	CCA	AGA	CCT	CAC Val	GAA	CIT	CCA	Thr	TYES
Gly Thr	Ser Asp	rys Asp	ASII	GLY	Jer	O.J	102	DÇU	01 0	013	****	D , 0.
	250		260		_	270			21	80		
• • • • • • • • • • • • • • • • • • •	***	AAA GCA	444	ፈ ጉጉ	ACA	TTA	GCT	GAC	GAT	CTA	AGT	AAA
CTA CTG	TTT TC	TTT CG1	TTT	AAT	TGT	TAA	CGA	CTG	CTA	GAT	TCA	TTT
Asp Asp	·Lys Ser	Lys Ala	Lys	Leu	Thr	Ile	Ala	Asp	qaA .	Leu	Ser	Lys>
200	300		31				320			330		
290 •	300			•	•		•		•	•		•
ACC ACA	TTC GA	CTT TT	AAA A	GAA	GAT	GGC	AAA	ACA	TTA	GTG	TCA	AGA
TGG TGT	AAG CT	GAA AA	TTT	CTT	CTA	CCG	TIT	TGI	AAT	CAC	AGT	ACT
Thr Thr	Phe Gl	Leu Le	ı Lys	Glu	Asp	GIY	Lys	Thr	Leu	vai	Ser	ALG.
340		350		360			3	70			380	
•	•	•	•	•		•		•	•		•	
AAA GTA	AGT TC	T AGA GA	C AAA	ACA	TCA	ACA ACA	GAT	GAA	OTA A	TTC	raa :	CTT
TTT CAT	TCA AG	A TCT CT r Arg As	o TTT	TGT	AG1	161 The	· CTA	((1)	TAC	, AAC Pha	, TIL	Glus
Lys Val	ser se	r wid wa	h ras	IIII	261		. waf	. 516	13C C	. T.11C		

FIGURE 24 (1 of 3)

B-31 OSP A/ PGAU OSP A FUSION

390	4	00	410	42	:0	430
•	•	• •	•	•		•
AAA GGT G	A TTG TCT	GCY YYY	ACC ATG	ACA AGA GA	LA AAT GGA	ACC AAA
TTT CCA CT	M AAC AGA	CGT TTT	TGG TAC	TGT TCT CT	T TTA CCT	TGG TTT
Lys Gly G	lu Leu Ser	Ala Lys	Thr Met	Thr Arg Gl	u Asn Gly	Thr Lys>
. 440)	450	46	50	470	460
•	• •	•	•	•	•	• •
CTT GAA T	AT ACA GAP	AAA DTA	AGC GAT	GGA ACC GG	SA AAA GCT	aaa gaa
GAA CTT A'	TA TGT CTT	TAC TTT	TCG CTA	CCT TGG CC	T TTT CGA	TIT CIT
Leu Glu T	yr Thr Glu	Met Lys	Ser Asp	Gly Thr Gl	ly Lys Ala	Lys Glu>
	•	500		510	520	
	490	500	•	• •	•	•
		·	CAA GGA	AAA GTA GO	T AAT GAT	AAA GTA
GTT TTA A	AA AAG 117	TCD CDD	CTT CCT	TTT CAT CO	SA PTA CTA	TTT CAT
CAA AAT 1	II IIC AA	Thr Leu	Glu Glv	Lys Val Al	la Asn Asp	Lvs Val>
Val Leu L	ys Lys Fin	: IIII Deu	010 017		· · · · · · · · · · · · · · · · · ·	
530	540	5	50	560	570	
• •	•	•	. • •	•	• •	•
ACA TTG G	AA GTA AA	A GAA GGA	ACC GTT	ACT TTA AC	GT AAG GAA	ATT GCA
TGT AAC C	TT CAT TT	r ctt cct	TGG CAA	TGA AAT TO	CA TTC CTT	TAA CGT
Thr Leu G	lu Val Ly	s Glu Gly	Thr Val	Thr Leu Se	er Lys Glu	ile Ala>
580	590		600	610		620
•	• •	•	•	•	•	
AAA TCT C	GA GAA GT	A ACA GTI	GCT CTT	AAT GAC A	CT AAC ACT	ACT CAG
TTT AGA C	CT CTT CA	T TGT CAP	CGA GAA	TTA CTG	GA TTG TGA	TGA GIC
Lys Ser (Gly Glu Va	1 Thr Val	Ala Leu	Asn Asp T	nr Asn Inr	TER GIES
630		640	650	. 6	60	670
• •	•	•	•	•	•	•
GCT ACT	DA AAA AAA	T GGC GC	A TGG GAT	TCA AAA A	CT TCT ACT	TIA ACA
CGA TGA	TTT TTT TO	A CCG CG	r acc cta	AGT TTT T	GA AGA TGA	AAT TG2
Ala Thr	Lys Lys Th	r Gly Ala	a Trp Asp	Ser Lys T	hr Ser Thr	Leu Into
6	80	690	•	700	710	720
•	•	•	•	• •	•	
ATT AGT	GTT AAC AG	C AAA AA	A ACT AC	A CAA CTT C	TG TTT ACT	AAA CAA
TAA TCA	CAA TTG T	CG TTT TT	T TGA TG	T GTT GAA	CAC AAA TG	A TTT GIT
Ile Ser	Val Asn S	er Lys Ly	s Thr Th	r Gln Leu V	val Phe Thi	r Lys Gln>
	730	740		750	760	
•	•		•	•	•	•
TAC ACA	ATA ACT G	TA AAA CA	A TAC GA	C TCC GCA	GGT ACC AA	T TTA GAA
ATG TGT	TAT TGA C	AT TTT GI	T ATG CI	G AGG CGT	CCA TGG TT	A AAT CIT
Tyr Thr	Ile Thr V	al Lys G	in Tyr As	p Ser Ala	Gly Thr As	n Leu Glu>
-						

FIGURE 24 (2 of 3)

B-31 OSP A /PGAU OSP A FUSION

770 780 790 800 810

GGC ACA GCA GTC GAA ATT AAA ACA CTT GAT GAA CTT AAA AAC GCT TTA

CCG TGT CGT CAG CTT TAA TTT TGT GAA CTA CTT GAA TTT TTG CGA AAT

Gly Thr Ala Val Glu Ile Lys Thr Leu Asp Glu Leu Lys Asn Ala Leu>

820

AAA TAA TTT ATT Lys ***>

B31/K48 fusion

20 10 ATG AAA AAA TAT TTA TTG GGA ATA GGT CTA ATA TTA GCC TTA ATA GCA TAC TIT TIT ATA AAT AAC CCT TAT CCA GAT TAT AAT CGG AAT TAT CGT Met Lys Lys Tyr Leu Leu Gly Ile Gly Leu Ile Leu Ala Leu Ile Ala> 90 80 70 50 60 TGT AAG CAA AAT GTT AGC AGC CTT GAC GAG AAA AAC AGC GTT TCA GTA ACA TTC GTT TTA CAA TCG TCG GAA CTG CTC TTT TTG TCG CAA AGT CAT Cys Lys Gln Asn Val Ser Ser Leu Asp Glu Lys Asn Ser Val Ser Val> 130 140 120 110 100 GAT TTG CCT GGT GAA ATG AAA GTT CTT GTA AGC AAA GAA AAA AAC AAA CTA AAC GGA CCA CTT TAC TTT CAA GAA CAT TCG TTT CTT TTT TTG TTT Asp Leu Pro Gly Glu Met Lys Val Leu Val Ser Lys Glu Lys Asn Lys> 180 170 160 150 GAC GGC AAG TAC GAT CTA ATT GCA ACA GTA GAC AAG CTT GAG CTT AAA CTG CCG TTC ATG CTA GAT TAA CGT TGT CAT CTG TTC GAA CTC GAA TTT Asp Gly Lys Tyr Asp Leu Ile Ala Thr Val Asp Lys Leu Glu Leu Lys> 220 230 210 200 GGA ACT TCT GAT AAA AAC AAT GGA TCT GGA GTA CTT GAA GGC GTA AAA CCT TGA AGA CTA TTT TTG TTA CCT AGA CCT CAT GAA CTT CCG CAT TTT Gly Thr Ser Asp Lys Asn Asn Gly Ser Gly Val Leu Glu Gly Val Lys> 260 270 280 250 GCT GAC AAA AGT AAA GTA AAA TTA ACA ATT TCT GAC GAT CTA GGT CAA CGA CTG TTT TCA TTT CAT TTT AAT TGT TAA AGA CTG CTA GAT CCA GTT Ala Asp Lys Ser Lys Val Lys Leu Thr Ile Ser Asp Asp Leu Gly Gln> 320 330 310 290 300 ACC ACA CTT GAA GTT TTC AAA GAA GAT GGC AAA ACA CTA GTA TCA AAA TGG TGT GAA CTT CAA AAG TTT CTT CTA CCG TTT TGT GAT CAT AGT TTT Thr Thr Leu Glu Val Phe Lys Glu Asp Gly Lys Thr Leu Val Ser Lys> 370 350 360 340 AAA GTA ACT TCC AAA GAC AAG TCA TCA ACA GAA GAA AAA TTC AAT GAA TTT CAT TGA AGG TTT CTG TTC AGT AGT TGT CTT CTT TTT AAG TTA CTT Lys Val Thr Ser Lys Asp Lys Ser Ser Thr Glu Glu Lys Phe Asn Glu> 420 410 400 390 AAA GGT GAA GTA TCT GAA AAA ATA ATA ACA AGA GCA GAC GGA ACC AGA TIT CCA CIT CAT AGA CIT TIT TAT TAT TGT TCT CGT CTG CCT TGG TCT Lys Gly Glu Val Ser Glu Lys Ile Ile Thr Arg Ala Asp Gly Thr Arg>

FIGURE 25 (1 of 2)

B31/K48 fusion

450 CTT GAA TAC ACA GGA ATT AAA AGC GAT GGA TCT GGA AAA GCT AAA GAG GAA CTT ATG TGT CCT TAA TTT TCG CTA CCT AGA CCT TTT CGA TTT CTC Leu Glu Tyr Thr Gly Ile Lys Ser Asp Gly Ser Gly Lys Ala Lys Glu> 500 510 490 520 GTT TTA AAA GGC TAT GTT CTT GAA GGA ACT CTA ACT GCT GAA AAA ACA CAA AAT TIT CCG ATA CAA GAA CTT CCT TGA GAT TGA CGA CTT TIT TGT Val Leu Lys Gly Tyr Val Leu Glu Gly Thr Leu Thr Ala Glu Lys Thr> : `=:::: 550 560 530 540 ACA TTG GTG GTT AAA GAA GGA ACT GTT ACT TTA AGC AAA AAT ATT TCA TGT AAC CAC CAA TTT CTT CCT TGA CAA TGA AAT TCG TTT TTA TAA AGT Thr Leu Val Val Lys Glu Gly Thr Val Thr Leu Ser Lys Asn Ile Ser> 600 610 580 590 AAA TCT GGG GAA GTT TCA GTT GAA CTT AAT GAC ACT GAC AGT AGT GCT TTT AGA CCC CTT CAA AGT CAA CTT GAA TTA CTG TGA CTG TCA TCA CGA Lys Ser Gly Glu Val Ser Val Glu Leu Asn Asp Thr Asp Ser Ser Ala> 630 640 650 660 670 GCT ACT AAA AAA ACT GCA GCT TGG AAT TCA AAA ACT TCC ACT TTA ACA CGA TGA TTT TTT TGA CGT CGA ACC TTA AGT TTT TGA AGG TGA AAT TGT Ala Thr Lys Lys Thr Ala Ala Trp Asn Ser Lys Thr Ser Thr Leu Thr> 680 690 700 710 720 ATT AGT GTG AAT AGC CAA AAA ACC AAA AAC CTT GTA TTC ACA AAA GAA TAA TCA CAC TTA TCG GTT TTT TGG TTT TTG GAA CAT AAG TGT TTT CTT Ile Ser Val Asn Ser Gln Lys Thr Lys Asn Leu Val Phe Thr Lys Glu> 740 750 760 GAC ACA ATA ACA GTA CAA AAA TAC GAC TCA GCA GGC ACC AAT CTA GAA CTG TGT TAT TGT CAT GTT TTT ATG CTG AGT CGT CCG TGG TTA GAT CTT Asp Thr Ile Thr Val Gln Lys Tyr Asp Ser Ala Gly Thr Asn Leu Glu> 770 780 800 810 GGC AAA GCA GTC GAA ATT ACA ACA CTT AAA GAA CTT AAA AAC GCT TTA CCG TTT CGT CAG CTT TAA TGT TGT GAA TTT CTT GAA TTT TTG CGA AAT

820

AAA TAA TTT ATT Lys ***>

Gly Lys Ala Val Glu Ile Thr Thr Leu Lys Glu Leu Lys Asn Ala Leu>

B-31 OSP A 25015 OSP A FUSION

	•	_	_			20			30			4	0			
		3	.0			20			•		•		•	•		
	•		TAT	-	~~~	CCN	ara	CGT	СТА	ATA	ATT	GCC	TTA	ATA	GCA	
ATG	AAA	AAA	ATA	TIM	770	CCT	TAT	CCA	GAT	TAT	AAT	CGG	TAA	TAT	CGT	
TAC	TIT	TIT	TIA	Len	Leu	Glv	Ile	Gly	Leu	Ile	Leu	Ala	Leu	Ile	Ala>	
met	Lys	Lys	TYT	Dea		,		-								
50			60			•	70			80			90			
		•	•		•		•	. •		•		•	•		•	
TGT	AAG	CAA	TAA	GTT	AGC	AGC	CTT	GAC	GAG	AAA	AAC	AGC	GTT	TCA	GTA	
	****	CTT	ጥጥ እ	CAA	TCG	TCG	GAA	CTG	CIC	1-1-L	116	160	CAA	YO1.	CAT	
Cys	Lys	Gln	Asn	Val	Ser	Ser	Leu	Asp	GIU	ĻYS.	'WZII	ser	val	261	Val>	
							120			1:	30		1	140		
10	00			110		•	120	-	•		•	•	-	•		
~~~	4.	CCT	GGT	CAA	ATG	AAA	GTT	CTT	GTA	AGC	AAA	GAA	AAA	AAC	AAA	
CTA.		CCA	CCA	CTT	TAC	TTT	CAA	GAA	CAT	TCG	TIT	CIT	TTT	TTG	TIT	
ASD	Leu	Pro	Gly	Glu	Met	Lys	Val	Leu	Val	Ser	Lys	G1u	Lys	Asn	Lys>	
1.Jp			-									•				
	150			1	60			170		_	180			1	90	
•	•		•		•	•			CT.	CAC		СТТ	CNC	حث.		
GAC	GGC	AAG	TAC	GAT	CTA	ATT	GCA	NCA TOT	CIA	CAC	TTC	GAA	CTC	GAA	AAA TTT	
CIG	CCG	TTC	ATG	CIA	LGAI	177	Ala	Thr	· Val	Asp	Lys	Leu	Glu	Leu	Lys>	
Asp	GIY	Lys	Tyr	YSL	, Dec					•	-				_	
		200			210	)		. 2	20			230			240	
•				•	•	•	•		•	•		•		•	•	
GGA	, ACI	r TC	r GAT	KAA 1	AA A	CAA	r GG	TCI	GGZ	\ GTA	CM	` GAA	GGC	GTA	AAA	
CCI	TG	A AG	A CT	A TT	r TT	TT	A CC	r AGA	CCI	r CAl	CAA	CIT		CAI	TTT	
Gly	Thi	r Se	r Ası	p Ly:	s Ası	n Ası	i GI	y Sei	. 617	y vai	Let	GIC	GIY	447	Lys>	
			250			260			27	0		2	280			
			250		•	•		•		•	•		•	•	r	
GC	r GA	C AA	A AG	AA T	A GT	A AA	A TT	A AC	TA A	T TC	r GAG	C GAT	r cta	GG7	KAD 1	
		~ ~~	ጥ ጥር	A TT	T CA	T TT	T AA	T TG	T TA	A AG	A CTO	G CIV	A GAT		4 GII	
Al	a As	p Ly	s Se	r Ly	s Va	l Ly	s Le	u Th	r Il	e Sei	r Asj	p As	o Lei	1 (17)	y Gln>	
				_			210			320			330	n		
290		_	30	0 .			310		•	320 *		•		•	•	
•	a .c		~r ca	A GT	T T1	'C AA	A GA	A GA	T GG	C AA	A AC	A CT	A GT	A TC	<b>AAA</b> A	
- m-	~ ~~	T (2)	<b>A</b>	ጥ ሮጀ	A A	G TI	TCI	TCI	'A CC	G TT	T 1G	T. CW	TCA	1 MG	1 11-	
Th	r Th	ir Le	eu Gl	lu Va	al Ph	ie Ly	s G1	lu As	p_G1	y Ly	s Th	r Le	u Va	1 Se	r Lys>	
***	- ••	,									•					
	340			350	)	_	3 (	50			370			380	! :	
	•		•		•	•		• ~ ~ ~	, ,, ,,	- ግክ መቅ	ים או מים	A A A	, T-1		T GAA	
A	AA G	A AT	CT TO	CC A	AA G	AC A	RG IV	CA TO	ጉሉ ብረ የጥ ጥር	CT (**	ייי פי	ות עו מי ייי	T A	G TI	AT GAA PA CTT	
77	rr c	T TA	GA A	GG T	TT C	so t	nc c	er S	er Ti	hr Gl	lu Gl	lu L	s Pt	ie As	sn Glu	>
L	ys V	al T	nr S	GI D	ys A	ah n	, , ,	- J	*							

FIGURE 26 (1 of 3)

#### B-31 OSP A/ 25015 OSP A FUSION

	390			40	00			410			420			4	30
•	•		•		•	•		•		•	•		•		•
AAA	GGT	GAA	GTA	TCT	GAA	AAA	ATA	ATA	ACA	AGA	GCA	GAC	GGA	ACC	λGA
TTT	CCA	CTT	CAT	AGA	CTT	TTT	TAT	TAT	TGT	TCT	CGT	CTG	CCT	TGG	TCT
Lys	GJA	Glu	Val	Ser	Glu	Lys	Ile	Ile	Thr	Arg	Ala	Аsp	Gly	Thr	Arg>
		140			450			. 4	60			470			400
·•		•		•	•		•	•	•	•		•		•	480
CTT	GAA	TAC	ACA	GGA	ATT	AAA	AGC	GAT	GGA	TCT	GGA	AAA	GCT	AAA	GAG
GAA	CTT	ATG	TGT	CCT	TAA	TTT	TCG	CTA	CCT	AGA	CCT	TTT	CGA	TATE	CTC
Leu	Glu	Tyr	Thr	Gly	Ile	Lys	Ser	Asp	Glỳ	Ser	Gly	Lys	Ala	Lys	Glu>
		49	90		9	500			510			5:	20		
	•		•	•		•		•	•		*		• •	٠.	
GTT	TTA	AAA	GGC	TAT	GTT	CTT	GAA	GGA	ACT	CTA	ACT	GCT	GAA	AAA	ACA
CAA	TAA	TTT	CCG	ATA	CAA	GAA	CTT	CCT	TGA	GAT	TGA	CGA	CTT	TTT	TGT
Val	Leu	Lys	Gly	Tyr	Val	Leu	Glu	Gly	Thr	Leu	Thr	Ala	Glu	Lys	Thr>
530			540			55	50		9	560			570		
•		•	•		•		•	•		•		•	•		•
ACA	TTG	GTG	GTT	AAA	GAA	GGA	ACT	GTT	ACT	ATT	AGC	AAA	TAA	TTA	TCA
TGT	AAC	CAC	CAA	TTT	CTT	CCT	TGA	CAA	TGA	TAA	TCG	TTT	ATT	TAA	AGT
Thr	Leu	Val	Val	Lys	Glu	Gly	Thr	Val	Thr	Leu	Ser	Lys	Asn	Ile	Ser>
58	30		-	590			600			61	.0		•	520	
	•	•		•		•	•		•		•	•		•	
AAA	· TCT	GGG	GAA	• GTT	TCA	GTT	GAA	CTT	AAT	GAC	ACT	GAC	AGT	AGT	GCT
AAA TTT	TCT AGA	CCC	GAA CTT	GTT CAA	AGT	CAA	GAA CTT	GAA	TTA	GAC CTG	ACT TGA	CTG	AGT TCA	AGT TCA	CGA
AAA TTT	TCT AGA	CCC	GAA CTT	GTT CAA	AGT	CAA	GAA CTT	GAA	TTA	GAC CTG	- ACT	CTG	AGT TCA	AGT TCA	CGA
AAA TTT	TCT AGA	CCC	GAA CTT	GTT CAA	AGT Ser	CAA	GAA CTT Glu	GAA	TTA	GAC CTG	ACT TGA	CTG	AGT TCA	AGT TCA	CGA Ala>
AAA TTT Lys	TCT AGA Ser 630	CCC	GAA CTT Glu	GTT CAA Val	AGT Ser	CAA Val	GAA CTT Glu	GAA Leu 550	TTA Asn	GAC CTG Asp	ACT TGA Thr	CTG Asp	AGT TCA Ser	AGT TCA Ser	CGA Ala>
AAA TTT Lys • GCT	TCT AGA Ser 630	CCC Gly AAA	GAA CTT Glu	GTT CAA Val	AGT Ser 10 • GCA	CAA Val	GAA CTT Glu	GAA Leu 550 •	TTA Asn GCA	GAC CTG Asp	ACT TGA Thr 660	CTG Asp	AGT TCA Ser	AGT TCA Ser 67	CGA Ala>
AAA TTT Lys GCT CGA	TCT AGA Ser 630 ACT TGA	CCC Gly AAA TTT	GAA CTT Glu • AAA TTT	GTT CAA Val 64 ACT TGA	AGT Ser 10 GCA CGT	CAA Val GCT CGA	GAA CTT Glu TGG ACC	GAA Leu 550 AAT TTA	TTA Asn GCA CGT	GAC CTG Asp	ACT TGA Thr 660 ACT	CTG Asp TCA AGT	AGT TCA Ser • ACT TGA	AGT TCA Ser 67 TTA AAT	CGA Ala> C ACA TGT
AAA TTT Lys GCT CGA	TCT AGA Ser 630 ACT TGA	CCC Gly AAA TTT	GAA CTT Glu • AAA TTT	GTT CAA Val 64 ACT TGA	AGT Ser 10 GCA CGT	CAA Val GCT CGA	GAA CTT Glu TGG ACC	GAA Leu 550 AAT TTA	TTA Asn GCA CGT	GAC CTG Asp	ACT TGA Thr 660	CTG Asp TCA AGT	AGT TCA Ser • ACT TGA	AGT TCA Ser 67 TTA AAT	CGA Ala> C ACA TGT
AAA TTT Lys GCT CGA	TCT AGA Ser 630 ACT TGA Thr	CCC Gly AAA TTT	GAA CTT Glu • AAA TTT	GTT CAA Val 64 ACT TGA	AGT Ser 10 GCA CGT	CAA Val GCT CGA	GAA CTT Glu TGG ACC	GAA Leu 550 * AAT TTA Asn	TTA Asn GCA CGT	GAC CTG Asp	ACT TGA Thr 660 ACT TGA Thr	CTG Asp TCA AGT	AGT TCA Ser • ACT TGA	AGT TCA Ser 67 TTA AAT	CGA Ala> C ACA TGT
AAA TTT Lys • GCT CGA Ala	TCT AGA Ser 630 ACT TGA Thr	CCC Gly AAA TTT Lys	GAA CTT Glu • AAA TTT Lys	GTT CAA Val 64 ACT TGA Thr	AGT Ser 10 GCA CGT Ala 690	CAA Val GCT CGA Ala	GAA CTT Glu TGG ACC Trp	GAA Leu 550 • AAT TTA Asn	TTA Asn GCA CGT Ala	GAC CTG Asp GGC CCG Gly	ACT TGA Thr 660 ACT TGA Thr	TCA AGT Ser	AGT TCA Ser • ACT TGA Thr	AGT TCA Ser 67 TTA AAT Leu	CGA Ala> C ACA TGT Thr> 720
AAA TTT Lys GCT CGA Ala	TCT AGA Ser 630 ACT TGA Thr	AAA TTT Lys	GAA CTT Glu AAA TTT Lys	GTT CAA Val 66 ACT TGA Thr	AGT Ser 10 GCA CGT Ala 690	CAA Val GCT CGA Ala	GAA CTT Glu TGG ACC Trp	GAA Leu 550 AAT TTA ASI 70	GCA CGT Ala	GAC CTG Asp GGC CCG Gly	ACT TGA Thr 660 ACT TGA Thr	TCA AGT Ser	AGT TCA Ser ACT TGA Thr	AGT TCA Ser 67 TTA AAT Leu	CGA Ala> C ACA TGT Thr> 720 CAA
AAA TTT Lys GCT CGA Ala	TCT AGA Ser 630 ACT TGA Thr ACT TGA	AAA TTT Lys 680 GTA CAT	GAA CTT Glu AAA TTT Lys	GTT CAA Val 64 ACT TGA Thr AAC	AGT Ser 10 GCA CGT Ala 690 AAA TTT	CAA Val GCT CGA Ala AAA TTT	GAA CTT Glu TGG ACC Trp	GAA Leu 550 AAT TTA ASN 70	GCA CGT Ala 00 GCC CGG	GAC CTG Asp GGC CCG Gly CTT GAA	ACT TGA Thr 660 ACT TGA Thr	TCA AGT Ser 110 TTT AAA	AGT TCA Ser ACT TGA Thr	AGT TCA Ser 67 TTA AAT Leu AAA TTT	CGA Ala> C ACA TGT Thr> 720 CAA GTT
AAA TTT Lys GCT CGA Ala	TCT AGA Ser 630 ACT TGA Thr ACT TGA	AAA TTT Lys 680 GTA CAT Val	GAA CTT Glu AAA TTT Lys AAC TTG ASn	GTT CAA Val 64 ACT TGA Thr AAC	AGT Ser 10 GCA CGT Ala 690 AAA TTT	CAA Val GCT CGA Ala AAA TTT	GAA CTT Glu TGG ACC Trp	GAA Leu 550 AAT TTA ASN 70	GCA CGT Ala 00 GCC CGG	GAC CTG Asp GGC CCG Gly CTT GAA	ACT TGA Thr 660 ACT TGA Thr	TCA AGT Ser 110 TTT AAA	AGT TCA Ser ACT TGA Thr	AGT TCA Ser 67 TTA AAT Leu AAA TTT	CGA Ala> C ACA TGT Thr> 720 CAA
AAA TTT Lys GCT CGA Ala	TCT AGA Ser 630 ACT TGA Thr ACT TGA	AAA TTT Lys 680 GTA CAT Val	GAA CTT Glu AAA TTT Lys	GTT CAA Val 64 ACT TGA Thr AAC	AGT Ser 10 GCA CGT Ala 690 AAA TTT Lys	CAA Val GCT CGA Ala AAA TTT	GAA CTT Glu TGG ACC Trp	GAA Leu 550 AAT TTA ASN 70	GCA CGT Ala 00 GCC CGG	GAC CTG Asp GGC CCG Gly CTT GAA	ACT TGA Thr 660 ACT TGA Thr	TCA AGT Ser 110 TTT AAA	AGT TCA Ser ACT TGA Thr ACA TGT Thr	AGT TCA Ser 67 TTA AAT Leu AAA TTT	CGA Ala> C ACA TGT Thr> 720 CAA GTT
AAA TTT Lys GCT CGA Ala ATT TAA Ile	TCT AGA Ser 630 ACT TGA Thr ACT TGA Thr	AAA TTT Lys 680 GTA CAT Val	GAA CTT Glu AAA TTT Lys AAC TTG Asn	GTT CAA Val 64 ACT TGA Thr AAC TTG Asn	AGT Ser 10 GCA CGT Ala 690 AAA TTT Lys	CAA Val GCT CGA Ala AAA TTT Lys	GAA CTT Glu TGG ACC Trp ACT TGA Thr	GAA Leu 550 AAT TTA ASN 70 AAA TTT Lys	GCA CGT Ala O GCC CGG Ala 750	GAC CTG Asp GGC CCG Gly CTT GAA Leu	ACT TGA Thr 660 ACT TGA Thr GTA CAT Val	TCA AGT Ser 710 TTT AAA Phe	AGT TCA Ser ACT TGA Thr ACA TGT Thr	AGT TCA Ser 67 TTA AAT Leu AAA TTT Lys	CGA Ala> C ACA TGT Thr> 720 CAA GTT Gln>
AAA TTT Lys GCT CGA Ala ATT TAA Ile	TCT AGA Ser 630 ACT TGA Thr ACT TGA Thr	CCC Gly AAA TTT Lys 680 GTA CAT Val	GAA CTT Glu AAA TTT Lys AAC TTG Asn	GTT CAA Val 64 ACT TGA Thr AAC TTG Asn	AGT Ser 10 GCA CGT Ala 690 AAA TTT Lys	CAA Val GCT CGA Ala AAA TTT Lys	GAA CTT Glu TGG ACC Trp ACT TGA Thr	GAA Leu 550 AAT TTA ASN 70 AAA TTT Lys	GCA CGT Ala O GCC CGG Ala 750	GAC CTG Asp GGC CCG Gly CTT GAA Leu	ACT TGA Thr 660 ACT TGA Thr GTA CAT Val	TCA AGT Ser 710 TTT AAA Phe	AGT TCA Ser ACT TGA Thr ACA TGT Thr	AGT TCA Ser 67 TTA AAT Leu AAA TTT Lys	CGA Ala>  C ACA TGT Thr>  720 CAA GTT Gln>
AAA TTT Lys GCT CGA Ala ATT TAA Ile	TCT AGA Ser 630 ACT TGA Thr ACT TGA Thr ACT TGA Thr	CCC Gly AAA TTT Lys 680 GTA CAT Val ATT TAA	GAA CTT Glu AAA TTT Lys AAC TTG Asn	GTT CAA Val 64 ACT TGA Thr AAC TTG Asn TCA AGT	AGT Ser 10 GCA CGT Ala 690 AAA TTT Lys CAA GTT	CAA Val GCT CGA Ala AAA TTT Lys	GAA CTT Glu TGG ACC Trp ACT TGA Thr	GAA Leu 550 AAT TTA ASN 70 AAA TTT Lys	GCA CGT Ala OCC CGG Ala 750 TCA AGT	GAC CTG Asp GGC CCG Gly CTT GAA Leu GCA	ACT TGA Thr 660 ACT TGA Thr GTA CAT Val	TCA AGT Ser 710 TTT AAA Phe ACC TGG	AGT TCA Ser ACT TGA Thr ACA TGT Thr	AGT TCA Ser 67 TTA AAT Leu AAA TTT Lys TTG AAC	CGA Ala>  C ACA TGT Thr>  720 CAA GTT Gln>

FIGURE 26 (2 of 3)

49//33

# B-31 OSP A/ 25015 OSP A FUSION

770 780 790 800 810

GGC ACA GCA GTC GAA ATT AAA ACA CTT GAT GAA CTT AAA AAC GCT TTA

CCG TGT CGT CAG CTT TAA TTT TGT GAA CTA CTT GAA TTT TTG CGA AAT

Gly Thr Ala Val Glu Ile Lys Thr Leu Asp Glu Leu Lys Asn Ala Leu>

AGA TCT

Arg>

FIGURE 26 (3 of 3)

### K48 OSP A/ B-31 OSP A/ K48 OSP A FUSION

10	20	30	4	0
• •	• •	• •	•	•
ATG AAA AAA TAT	TTA TTG GGA	ATA GGT CTA	ATA TTA GCC	TTA ATA GCA
TAC TTT TTT ATA	AAT AAC CCT	TAT CCA GAT	TAT AAT CGG	AAT TAT CGT
Met Lys Lys Tyr	ren ren Già	ite Giy Leu	Tie Deu Ala	red lie Mia>
50 60	7	70	80	90
• •	•	- · · · ·		COM DO: 07:
TGT AAG CAA AAT	GTT AGC AGC	CIT GAT GAA	TTT TTN TCC	GIT TCA GTA
ACA TTC GTT TTA Cys Lys Gln Asn	CAA 1CG 1CG	Lau Asp Glu	Tys Ash Ser	Val Son Val
CAS TAS GIU YEU	val ser ser	rea wab ora	Dys Man der	Agt Set Agt>
100	110	120	130	- 140
• •	• •	• •	•	•
GAT TTA CCT GGT	GGA ATG ACA	GTT CTT GTA	AGT AAA GAA	AAA GAC AAA
CTA AAT GGA CCA	CCT TAC TGT	CAA GAA CAT	TCA TIT CIT	TIT CIG TIT
Asp Leu Pro Gly	Gly Met Thr	Val Leu val	ser Lys Glu	Lys Asp Lys>
150	160	170	180	190
• •	• •	•	• •	•
GAC GGT AAA TAC	AGT CTA GAG	GCA ACA GTA	GAC AAG CTT	GAG CTT AAA
CTG CCA TTT ATG	TCA GAT CTC	CGT TGT CAT	CTG TTC GAA	CTC GAA TTT
Asp Gly Lys Tyr	Ser Leu Glu	Ala Thr Val	Asp Lys Leu	Glu Leu Lys>
200	210	220	230	240
•	• •	• • •	101 000 011	CCT CIL III
GGA ACT TCT GAT	AAA AAC AAC	GGT TCT GGA	TOT CAR CTT	CCE CTT TTT
CCT TGA AGA CTA	The lighted	Cly Ser Gly	Thr Leu Glu	Gly Glu Lyss
Gly Thr Ser Asp	Dys Asii Asii	dry ser or,	504 014	01, 011 2,01
250	260	270	21	80
ACT GAC AAA AGT		ምምአ አር <u>ን</u> አሞሞ	CCT CAT CAC	CTA AGT CAA
TGA CTG TTT TCA	, AAA GIA AAA . TTT CLT TTT	AAT TGT TAA	CGA CTA CTG	GAT TCA GTT
Thr Asp Lys Ser	Lvs Val Lvs	Leu Thr Ile	Ala Asp Asp	Leu Ser Gln>
THE KSP SJS GG				
290 300	3	10	320	330
ACT AAA TTT GAI	ል ልተተ ተተር ልልል	GAA GAT GCC	AAA ACA TTA	GTA TCA AAA
TGA TTT AAA CT	TAA AAG TTI	CTT CTA CGG	TTT TGT AAT	CAT AGT TTT
Thr Lys Phe Gli	u Ile Phe Lys	Glu Asp Ala	Lys Thr Leu	Val Ser Lys>
	-			•
340	350	360	370	380
AAA GTA ACC CT	T 222 CAC 234	י דרא דרא ארי	GAA GAA AAA	TTC AAC GAA
TTT CAT TGG GA	1 ለለለ <i>GAC</i> ለለ(	TON TOL TOL C	CTT CTT TTT	AAG TTG CTT
Lys Val Thr Le	u Lvs Asp Lvs	s Ser Ser Thi	Glu Glu Lys	Phe Asn Glu>
DYS TUL 1416 DC				

FIGURE 27 (1 of 3)

51/133

# K48 OSP A/ B-31 OSP A/ K48 OSP A FUSION

390	400	.41	0	420	430
AAG GGT GAA TTC CCA CTT					
Lys Gly Glu	Thr Ser Glu	Lys Thr I	Te var ims	•••	Gly Thr Arg>
440	450 • •	•	460	470	• •
	ACA GAC ATA				
Leu Glu Tyr	Thr Asp Ile	Lys Ser A	sp dry ser	, .,.	
	90	500	510	•	0
	GAC TTT ACT				
Val Leu Lys	s Asp Phe Th	r Leu Glu	dia tur pe	T VIS VIS	Asp Gly Lys>
530	540	550	560 • • • • • • • • • • • • • • • • • • •	 	•
					AAG ATT TCA TTC TAA AGT Lys Ile Ser>
		600		610	620
580 •	590	• • •	CTT AAT GA	· C ACT GAC	AGT AGT GCT
					TCA TCA CGA Ser Ser Ala>
630	640		650	660	670
• •	• • • • • • • • • • • • • • • • • • •	• CA GCT TGG	AAT TCA A	AA ACT TC	ACT TTA ACA
					TGA AAT TGT Thr Leu Thr>
68		590	700	710	720
ATT AGT C	TG AAT AGC	AA AAA ACC	AAA AAC C	TT GTA TT	C ACA AAA GAA G TGT TTT CTT Le Thr Lys Glu>
TAA TCA (	Val Asn Ser (	Gln Lys Th	Lys Asn I	eu Val Ph	e Thr Lys Glu>
_	730	740	750 • •	•	760
GAC ACA CTG TGT Asp Thr	ATA ACA GTA TAT TGT CAT Ile Thr Val	CAA AAA TA GTT TTT AT Gln Lys Ty	C GAC TCA G CTG AGT r Asp Ser	GCA GGC AGC CGT CCG TG	CC AAT CTA GAA GG TTA GAT CTT hr Asn Leu Glu

FIGURE 27 (2 of 3)

K48 OSP A / B-31 OSP A/ K48 OSP A FUSION

770 780 790 800 810

GGC AAA GCA GTC GAA ATT ACA ACA CTT AAA GAA CTT AAA AAC GCT TTA CCG TTT CGT CAG CTT TAA TGT TGT GAA TTT CTT GAA TTT TTG CGA AAT Gly Lys Ala Val Glu Ile Thr Thr Leu Lys Glu Leu Lys Asn Ala Leu>

820

AAA TAA

TTT ATT

Lys ***>

FIGURE 27 (3 of 3)

# B-31 OSP A/K48 OSP A/ B-31 OSP A/ K48 OSP A FUSION

10		20	30		40	
• •	•	•	•	•	•	•
ATG AAA AAA T	AT TTA TTG	GGA ATA	GGT CTA	ATA TTA	GCC TTA	ATA GCA
TAC TTT TTT A	TA AAT AAC	CCT TAT	CCA GAT	TAA TAT	CGG AAT	TAT CGT
Met Lys Lys T	yr Leu Leu	Gly Ile	Gly Leu	Ile Leu	Ala Leu	Ile Ala>
		70		80	90	
50	60	70	•	•	• •	•
TGT AAG CAA A	AT GTT AGO	AGC CTT	GAT GAA	AAA AAT	AGC GTT	TCA GTA
ACA TTC GTT T	TA CAA TCG	TCG GAA	CTA CTT	ATT-FFF	TCG CAA	AGT CAT
Cys Lys Gln A	sn Val Ser	Ser Leu	Asp Glu	Lys Asn	Ser Val	Ser Val>
•,• •,•						
100	110	120		130	- 3	L40·
• •	•	• •	•	•	•	•
GAT TTA CCT G	GT GGA ATC	ACA GTT	CTT GTA	AGT AAA	GAA AAA	GAC AAA
CTA AAT GGA C	CA CCT TAC	TGT CAA	GAA CAT	TCA TIT	CIT TIT	CIG TIT
Asp Leu Pro G	ily Gly met	inr vai	Leu vai	Ser Lys	GIG LYS	Asp Lys>
150	160		170	180		190
* *		•	•	• •	` <b>.</b>	•
GAC GGT AAA T						
CTG CCA TTT A	TG TCA GAT	CTC CGT	TGT CAT	CTG TTC	GAA CTC	GAA TTT
Asp Gly Lys T	lyr Ser Lei	Glu Ala	Thr Val	Asp Lys	Leu Glu	Leu Lys>
	21/		220	-		240
200	210	, • •	220	•	.50	240
GGA ACT TCT C	)AA AAA T45	AAC GGT	TCT GGA	ACA CTT	GAA GGT	GAA AAA
CCT TGA AGA C	TA TIT TI	TTG CCA	AGA CCT	TGT GAA	CTT CCA	CTT TTT
Gly Thr Ser A	Asp Lys Asi	n Asn Gly	Ser Gly	Thr Leu	Glu Gly	Glu Lys>
	•					
250	0	260	270	_	280	_
•	• •	• • • • • • • • • • • • • • • • • • • •	ics som	רכית כיית	CAC CTA	NCT CNA
ACT GAC AAA A	AGT AAA GI.	ከ11 አሌሌ ይ ኮልል ጥጥ ከ	TOT TAA	CCA CTA	CTG GAT	TCA GTT
Thr Asp Lys	Ser Ivs Va	l Lvs Leu	Thr Ile	Ala Asp	Asp Leu	Ser Gln>
THE ASP DYS	502 575 12	,				
290	300	310		320	330	
• •	• •	•	•	•	• •	•
ACT AAA TTT						
TGA TTT AAA	CTT TAA AA	G TIT CIT	CTA CGG	TTT TGT	AAT CAT	AGT TIT
Thr Lys Phe	Glu lle Pr	e ras cu	1 ASP Ala	Lys Inc	ren var	Ser Lyse
<del>-</del>						
340		360	0	370		380
340	350	360		370	•	380
• •	350	•	• •	•	. AAA TTO	•
340  AAA GTA ACC TTT CAT TGG Lys Val Thr	350 • CTT AAA GA GAA TTT CT	C AAG TC C TTC AG	• • A TCA ACA T AGT TG7	GAA GAA	TTT AAC	AAC GAA

FIGURE 28 (1 of 3)

## B-31 OSP A/K48 OSP A/ B-31 OSP A/ K48 OSP A FUSION

	390			4	00			410			420			4	30
•	•		•		•	•		. •		•	•		•		•
AAG	GGT	GAA	ACA	TCT	GAA	AAA	ACA	ATA	GTA	AGA	GCA	TAA	GGA	ACC	AGA
TTC	CCA	CTT	TGT	AGA	CTT	TTT	TGT	TAT	CAT	TCT	CGT	TTA	CCT	TGG	TCT
Lys	Gly	Glu	Thr	Ser	Glu	Lys	Thr	Ile	Val	Arg	Ala	Asn	Gly	Thr	Arg>
		440			450			4	60			470			400
•		•		•	. •		•		•	•		•		•	480
CTT	GAA	TAC	ACA	GAC	ATA	AAA	AGC	GAT	GGA	TCC	GGA	AAA	GCT	AAA	GAA
GAA	CTT	ATG	TGT	CTG	TAT	TTT	TCG	CTA	CCT	AGG	CCT	TATE	CCA	TTT	
Leu	Glu	Tyr	Thr	Asp	Ile	Lys	Ser	Аsp	Gľy	Sēr'	gly	Lys	Ala	Lys	Glu>
		45	90		9	500	•		510			52	20		
	•		•	•		•		•	•		•		•	•	•
GTT	TTA	AAA	GAC	TTT	ACT	CTT	GA_A	GGA	ACT	CTA	GCT	GCT	GAC	GGC	AAA
CAA	AAT	TTT	CTG	AAA	TGA	GAA	CTT	CCT	TGA	GAT	CGA	CGA	CTG	CCG	TTT
Val	Leu	Lys	Asp	Phe	Thr	Leu	Glu	Gly	Thr	Leu	Ala	Ala	Asp	Gly	Lys>
530			540			5	50		9	560			570		
•		•	•		•		•	•		•		•	r		•
ACA	ACA	TTG	AAA	GTT	ACA	GAA	GGC	ACT	GTT	GTT	TTA	AGC	AAG	TTA	TCA
TGT	TGT	AAC	TTT	CAA	TGT	CTT	CCG	TGA	CAA	CAA	TAA	TCG	TTC	AAT	AGT
Thr	Thr	Leu	Lys	Val	Thr	Glu	Gly	Thr	Val	Val	Leu	Ser	Lys	Ile	Ser>
S	30		9	90			600			61	.0		€	20	
	•	•		•		•	•		•		•	•		•	
AAA	TCT	GGG	GAA	• GTT	TCA	GTT	GAA	CTT	* AAT	GAC	• ACT	GAC	AGT	* AGT	GCT
AAA TTT	TCT AGA	CCC	GAA CTT	GTT CAA	AGT	CAA	GAA CTT	GAA	TTA	GAC CTG	• ACT TÇA	CTG	AGT TCA	AGT TCA	CGA
AAA TTT	TCT AGA	CCC	GAA CTT	GTT CAA	AGT	CAA	GAA CTT	GAA	TTA	GAC CTG	• ACT TÇA	CTG	AGT TCA	AGT TCA	GCT CGA Ala>
AAA TTT	TCT AGA	CCC	GAA CTT	GTT CAA	AGT Ser	CAA	GAA CTT Glu	GAA	TTA	GAC CTG	• ACT TÇA	CTG	AGT TCA	AGT TCA	CGA Ala>
AAA TTT Lys	TCT AGA Ser 630	CCC	GAA CTT Glu	GTT CAA Val	AGT Ser	CAA Val	GAA CTT Glu	GAA Leu 50	TTA Asn	GAC CTG Asp	ACT TGA Thr	CTG Asp	AGT TCA Ser	AGT TCA Ser	CGA Ala>
AAA TTT Lys • GCT	TCT AGA Ser 630	CCC Gly	GAA CTT Glu	GTT CAA Val	AGT Ser 10 GCA	CAA Val	GAA CTT Glu	GAA Leu 50	TTA Asn TCA	GAC CTG Asp	ACT TGA Thr 660	CTG Asp	AGT TCA Ser	AGT TCA Ser 67	CGA Ala> 0
AAA TTT Lys • GCT CGA	TCT AGA Ser 630 ACT TGA	CCC Gly AAA TTT	GAA CTT Glu • AAA TTT	GTT CAA Val 64 ACT TGA	AGT Ser 10 GCA CGT	CAA Val GCT CGA	GAA CTT Glu TGG ACC	GAA Leu 50 AAT TTA	TTA Asn TCA AGT	GAC CTG Asp	ACT TGA Thr 660 ACT	CTG Asp TCC AGG	AGT TCA Ser • ACT	AGT TCA Ser 67 TTA AAT	CGA Ala> 0 • ACA
AAA TTT Lys • GCT CGA	TCT AGA Ser 630 ACT TGA	CCC Gly AAA TTT	GAA CTT Glu • AAA TTT	GTT CAA Val 64 ACT TGA	AGT Ser 10 GCA CGT	CAA Val GCT CGA	GAA CTT Glu TGG ACC	GAA Leu 50 AAT TTA	TTA Asn TCA AGT	GAC CTG Asp	ACT TGA Thr 660 ACT	CTG Asp TCC AGG	AGT TCA Ser • ACT	AGT TCA Ser 67 TTA AAT	CGA Ala> 0
AAA TTT Lys • GCT CGA	TCT AGA Ser 630 ACT TGA Thr	CCC Gly AAA TTT	GAA CTT Glu • AAA TTT	GTT CAA Val 64 ACT TGA	AGT Ser 10 GCA CGT	CAA Val GCT CGA	GAA CTT Glu TGG ACC	GAA Leu 50 AAT TTA	TTA Asn TCA AGT Ser	GAC CTG Asp	ACT TGA Thr 660 ACT TGA Thr	CTG Asp TCC AGG	AGT TCA Ser • ACT	AGT TCA Ser 67 TTA AAT	CGA Ala> 0 • ACA
AAA TTT Lys GCT CGA Ala	TCT AGA Ser 630 ACT TGA Thr	AAA TTT Lys	GAA CTT Glu • AAA TTT Lys	GTT CAA Val 64 ACT TGA Thr	AGT Ser 10 GCA CGT Ala 690	CAA Val GCT CGA Ala	GAA CTT Glu TGG ACC Trp	GAA Leu 50 AAT TTA Asn	TTA Asn TCA AGT Ser	GAC CTG Asp AAA TTT Lys	ACT TGA Thr 660 ACT TGA Thr	TCC AGG Ser	AGT TCA Ser • ACT TGA Thr	AGT TCA Ser 67 TTA AAT Leu	CGA Ala>  0 ACA TGT Thr>
AAA TTT Lys GCT CGA Ala	TCT AGA Ser 630 ACT TGA Thr	AAA TTT Lys	GAA CTT Glu AAA TTT Lys	GTT CAA Val 64 ACT TGA Thr	AGT Ser 10 GCA CGT Ala 690 CAA	CAA Val GCT CGA Ala	GAA CTT Glu TGG ACC Trp	GAA Leu 550 AAT TTA ASD 70	TCA AGT Ser	GAC CTG Asp AAA TTT Lys	ACT TGA Thr 660 ACT TGA Thr	TCC AGG Ser	AGT TCA Ser • ACT TGA Thr	AGT TCA Ser 67 TTA AAT Leu	CGA Ala>  0 ACA TGT Thr> 720 GAA
AAA TTT Lys GCT CGA Ala	TCT AGA Ser 630 ACT TGA Thr AGT TCA	AAA TTT Lys	GAA CTT Glu AAA TTT Lys	GTT CAA Val 64 ACT TGA Thr AGC TCG	AGT Ser 10 GCA CGT Ala 690 CAA GTT	CAA Val GCT CGA Ala AAA TTT	GAA CTT Glu TGG ACC Trp	GAA Leu 550 AAT TTA ASD 70	TTA AST TCA AGT Ser 00 AAC TTG	GAC CTG Asp AAA TTT Lys	ACT TGA Thr 660 ACT TGA Thr	TCC AGG Ser	AGT TCA Ser • ACT TGA Thr	AGT TCA Ser 67 TTA AAT Leu	CGA Ala>  0 ACA TGT Thr> 720 GAA CTT
AAA TTT Lys GCT CGA Ala	TCT AGA Ser 630 ACT TGA Thr AGT TCA	AAA TTT Lys	GAA CTT Glu AAA TTT Lys	GTT CAA Val 64 ACT TGA Thr AGC TCG	AGT Ser 10 GCA CGT Ala 690 CAA GTT	CAA Val GCT CGA Ala AAA TTT	GAA CTT Glu TGG ACC Trp	GAA Leu 550 AAT TTA ASD 70	TTA AST TCA AGT Ser 00 AAC TTG	GAC CTG Asp AAA TTT Lys	ACT TGA Thr 660 ACT TGA Thr	TCC AGG Ser	AGT TCA Ser • ACT TGA Thr	AGT TCA Ser 67 TTA AAT Leu	CGA Ala>  0 ACA TGT Thr> 720 GAA
AAA TTT Lys GCT CGA Ala	TCT AGA Ser 630 ACT TGA Thr AGT TCA	AAA TTT Lys	GAA CTT Glu AAA TTT Lys AAT TTA Asn	GTT CAA Val 64 ACT TGA Thr AGC TCG	AGT Ser 10 GCA CGT Ala 690 CAA GTT Gln	CAA Val GCT CGA Ala AAA TTT	GAA CTT Glu TGG ACC Trp	GAA Leu 550 AAT TTA ASD 70	TTA AST TCA AGT Ser 00 AAC TTG	GAC CTG Asp AAA TTT Lys	ACT TGA Thr 660 ACT TGA Thr	TCC AGG Ser	AGT TCA Ser ACT TGA Thr ACA TGT Thr	AGT TCA Ser 67 TTA AAT Leu	CGA Ala>  0 ACA TGT Thr>  720 GAA CTT
AAA TTT Lys GCT CGA Ala ATT TAA Ile	TCT AGA Ser 630 ACT TGA Thr AGT TCA Ser	AAA TTT Lys 680 GTG CAC Val	GAA CTT Glu AAA TTT Lys AAT TTA Asn	GTT CAA Val 64 ACT TGA Thr AGC TCG Ser	AGT Ser 10 GCA CGT Ala 690 CAA GTT Gln	CAA Val GCT CGA Ala AAA TTT Lys	GAA CTT Glu TGG ACC Trp ACC TGG Thr	GAA Leu 50 AAT TTA Asn 70 AAA TTT Lys	TTA Asn TCA AGT Ser 00 AAC TTG Asn 750	GAC CTG Asp AAA TTT Lys CTT GAA Leu	ACT TGA Thr 660 ACT TGA Thr GTA CAT Val	TCC AGG Ser TTC AAG Phe	AGT TCA Ser ACT TGA Thr ACA TGT Thr	AGT TCA Ser 67 TTA AAT Leu AAA TTT Lys	CGA Ala>  0 ACA TGT Thr> 720 GAA CTT Glu>
AAA TTT Lys GCT CGA Ala ATT TAA Ile	TCT AGA Ser 630 ACT TGA Thr AGT TCA Ser	AAA TTT Lys 680 GTG CAC Val	GAA CTT Glu AAA TTT Lys AAT TTA Asn	GTT CAA Val 64 ACT TGA Thr AGC TCG Ser GTA	AGT Ser 10 GCA CGT Ala 690 CAA GTT Gln	CAA Val GCT CGA Ala AAA TTT Lys	GAA CTT Glu TGG ACC Trp ACC TGG Thr	GAA Leu 50 AAT TTA ASN 70 AAA TTT Lys	TTA Asn TCA AGT Ser 00 AAC TTG Asn 750	GAC CTG Asp AAA TTT Lys CTT GAA Leu	ACT TGA Thr 660 ACT TGA Thr GTA CAT Val	TCC AGG Ser TTC AAG Phe 76	AGT TCA Ser ACT TGA Thr ACA TGT Thr	AGT TCA Ser 67 TTA AAT Leu AAA TTT Lys	CGA Ala>  0  ACA TGT Thr> 720  GAA CTT Glu>
AAA TTT Lys GCT CGA Ala ATT TAA Ile	TCT AGA Ser 630 ACT TGA Thr AGT TCA Ser	AAA TTT Lys 680 CAC Val 73	GAA CTT Glu AAA TTT Lys AAT TTA ASD	GTT CAA Val 64 ACT TGA Thr AGC TCG Ser GTA CAT	AGT Ser 10 GCA CGT Ala 690 CAA GTT Gln	CAA Val GCT CGA Ala TTT Lys	GAA CTT Glu TGG ACC Trp ACC TGG Thr	GAA Leu 550 AAT TTA ASN 70 AAA TTT Lys	TCA AGT Ser O. AAC TTG ASN 750 TCA AGT	GAC CTG Asp AAA TTT Lys CTT GAA Leu GCA	ACT TGA Thr 660 ACT TGA Thr GTA CAT Val	TCC AGG Ser TTC AAG Phe ACC TGG	AGT TCA Ser ACT TGA Thr ACA TGT Thr	AGT TCA Ser 67 TTA AAT Leu AAA TTT Lys CTA GAT	CGA Ala>  0  ACA TGT Thr> 720  GAA CTT Glu>

FIGURE 28 (2 of 3)

## B-31 OSP A/K48 OSP A/ B-31 OSP A/ K48 OSP A FUSION

770 780 790 800 810

GGC AAA GCA GTC GAA ATT ACA ACA CTT AAA GAA CTT AAA AAC GCT TTA

CCG TTT CGT CAG CTT TAA TGT TGT GAA TTT CTT GAA TTT TTG CGA AAT

Gly Lys Ala Val Glu Ile Thr Thr Leu Lys Glu Leu Lys Asn Ala Leu>

820

AAA TAA TTT ATT Lys ***>

#### B-31 OSPA/ B-31 OSPB FUSION

20 ATG AAA AAA TAT ITA TTG GGA ATA GGT CTA ATA TTA GCC TTA ATA GCA TAC TIT TIT ATA AAT AAC CCT TAT CCA GAT TAT AAT CGG AAT TAT CGT Met Lys Lys Tyr Leu Leu Gly Ile Gly Leu Ile Leu Ala Leu Ile Ala> 90 70 80 60 50 TGT AAG CAA AAT GTT AGC AGC CTT GAC GAG AAA AAC AGC GTT TCA GTA ACA TTC GTT TTA CAA TCG TCG GAA CTG CTC TTT_TTG TCG CAA AGT CAT Cys Lys Gln Asn Val Ser Ser Leu Asp Glu Lys Asn Ser Val Ser Val> 130 120 110 GAT TTG CCT GGT GAA ATG AAA GTT CTT GTA AGC AAA GAA AAA AAC AAA CTA AAC GGA CCA CTT TAC TTT CAA GAA CAT TCG TTT CTT TTT TTG TTT Asp Leu Pro Gly Glu Met Lys Val Leu Val Ser Lys Glu Lys Asn Lys> 180 170 160 150 GAC GGC AAG TAC GAT CTA ATT GCA ACA GTA GAC AAG CTT GAG CTT AAA CTG CCG TTC ATG CTA GAT TAA CGT TGT CAT CTG TTC GAA CTC GAA TTT Asp Gly Lys Tyr Asp Leu Ile Ala Thr Val Asp Lys Leu Glu Leu Lys> 240 230 220 200 210 GGA ACT TCT GAT AAA AAC AAT GGA TCT GGA GTA CTT GAA GGC GTA AAA CCT TGA AGA CTA TTT TTG TTA CCT AGA CCT CAT GAA CTT CCG CAT TTT Gly Thr Ser Asp Lys Asn Asn Gly Ser Gly Val Leu Glu Gly Val Lys> 270 280 260 250 GCT GAC AAA AGT AAA GTA AAA TTA ACA ATT TCT GAC GAT CTA GGT CAA CGA CTG TTT TCA TTT CAT TTT AAT TGT TAA AGA CTG CTA GAT CCA GTT Ala Asp Lys Ser Lys Val Lys Leu Thr Ile Ser Asp Asp Leu Gly Gln> 330 320 310 300 ACC ACA CTT GAA GTT TTC AAA GAA GAT GGC AAA ACA CTA GTA TCA AAA TGG TGT GAA CTT CAA AAG TTT CTT CTA CCG TTT TGT GAT CAT AGT TTT Thr Thr Leu Glu Val Phe Lys Glu Asp Gly Lys Thr Leu Val Ser Lys> 370 360 350 340 AAA GTA ACT TCC AAA GAC AAG TCA TCA ACA GAA GAA AAA TTC AAT GAA TTT CAT TGA AGG TTT CTG TTC AGT AGT TGT CTT CTT TTT AAG TTA CTT Lys Val Thr Ser Lys Asp Lys Ser Ser Thr Glu Glu Lys Phe Asn Glu>

FIGURE 29 (1 of 3)

## B-31 OSP A/ B-31 OSP B FUSION

390	41	00	4:	10		4	20			430	)
	•	•		•	•		•		•	•	•
AAA GGT GAA	GTA TCT	GAA AAA	ATA .	ATA A	CA A	GA G	CA G	AC G	GA A	ACC 2	AGA
	C	$-\tau$	' TAT '	IAI I	.61 1	CI C	GI C	10 6		. نائ	101
Lys Gly Glu	Val Ser	Glu Lys	Ile	Ile T	thr A	rg A	la A	sp G	ily :	Thr A	krg>
ביים כבי								_			
440		450		460	)		47	U •		•	480
• •	•	•	•				מ מים			- ***	C N C
CTT GAA TAC	ACA GGA	AAA TTA	AGC	GAT	OW I	CIG	ሌለ ለ ውጥ ጥ	77 C	C N	ا تعلم العمري	CTC
GAA CTT ATG	TGT CCT	TAA TII	TCG	CIA (	51 5	SON C	111 1	11 C	la i	lie i	C1U
GAA CTT ATG Leu Glu Tyr	Thr Gly	Ile Lys	ser	ASP (	ıy s	er G	TY L	ys r	ila .	Lys	GIU>
	00	500		9	510			520	٥.		
4	90			•	•		•	•	•	•	
GTT TTA AAA	- 	י כדד כדי	r GAA	GGA 2	ACT (	TA AT	CT G	CT (	GAA .	AA.	ACA
	. CCC NTN	CAR CAR	TT	CCT	IGA C	JAI I	LGA C	.UK I	-11	41:	101
Val Leu Lys	יינת אור	· Val Le	ı Glu	Gly '	Thr I	Leu 1	thr A	la (	Glu	Lys	Thr>
Val Leu Lys	GIY IYI	. 461 50.									
530	540	!	550		56	60		:	570		
	•	•	•	•		•	•	•	•		•
ACA TTG GTG	GTT AN	A GAA GG	A ACT	GTT	ACT '	TTA 1	AGC A	AAA .	TAA	AT:	TCA
	~ ~ X X ~ ~ ~ ~	ד כשד ככ	T TGA	CAA	TGA A	AAT :	176 .	111	TIM	I V.	AG I
TGT AAC CAC	l Val Ly	s Glu Gl	y Thr	Val	Thr	Leu :	Ser 1	Lys	Asn	Ile	Ser>
200	•										
580	590		600		_	61	•		•	20	
•	• •	•		~~~		CNC	ት እርጥ	C N C	AGT	AGT.	CCT
AAA TCT GG	g gaa gt	T TCA GI	T GAA	CII	WAI	CTC .	TCD.	CTG	TCA	TCL	CGA
TTT AGA CC	C CTT CA	A AGT CA	A CIT	GAA	IIA	7.0	The	ACD	Sar	Ser	Alas
Lys Ser Gl	y Glu Va	l Ser Va	il Glu	Leu	ASII	vəh	1111	vəħ	561		
		640		65'0			660			€	70
630	•	•	•	•		•	•		•		•
GCT ACT AA	א מממ.	T GCA G	T TGG	TAA	GAC	AGT	ACT	AGC	ACT	TTA	λCλ
	~r ~r~r~r ~r	28 CCT C	RA ACC	TTA	CIG	TCA	TGA	100	IGN	WVI	101
Ala Thr Ly	's Lvs Ti	r Ala A	la Trp	Asn	Asp	Ser	Thr	Ser	Thr	Leu	Thr>
Ala IIII D	, <b>5</b>										
680	<b>)</b> .	690		7	00		-	710			720
•	•	•	•		•		~~~			, , ,	ርስጥ:
ATT AGT G	CT GAC A	GC AAA A	AA ACT	AAA 1	GAT	TTG	GIG	110	7.7%	י איניע י אינייע	CTA
	~ እ <i>~</i> ምላ	ע מארשה ט	77 71	4 111	CIM	NAC.	-	200	,,,,		
TAA TCA C	la Asp S	er Lys L	ys Th	r Lys	Asp	reu	val	rne	Det		. vob.
	-30	74	. ^		750	١		7	60		
_	730	•	+	•	•	•	•	•	•		•
GGT ACA A	- 	יים ארא הא	מד ממי	מא כ	. ACA	GCT	· GGA	ACC	AG	CT	A GAA
	** WCT (	እጥ ርጥጥ (	LA THE	ת אוני	3 TG1	' CGA	, CC1	100	, ,,,	G GY	
CCA TGT T Gly Thr I	AN IOI C	al Gln (	iln Tv	r Ası	n Thi	r Ala	Gly	Thi	Se	r Le	u Glu>
GIV TOT 1	'TG TIIT /	, ar arii ,					-				

FIGURE 29 (2 of 3)

B-31 OSP A/ B-31 OSP B FUSION

770 780 790 800 810

GGA TCA GCA AGT GAA ATT AAA AAT CTT TCA GAG CTT AAA AAC GCT TTA
CCT AGT CGT TCA CTT TAA TTT TTA GAA AGT CTC GAA TTT TTG CGA AAT

Gly Ser Ala Ser Glu Ile Lys Asn Leu Ser Glu Leu Lys Asn Ala Leu>

820

AAA TAA TTT ATT Lys ***>

٠,

59//33

#### B-31 OSP A/ B-31 OSP B / B-31 OSP C FUSION

nce Range: 1 to 1401

30 20 10 ATG AAA AAA TAT TTA TTG GGA ATA GGT CTA ATA TTA GCC TTA ATA GCA TAC TIT TIT ATA AAT AAC CCT TAT CCA GAT TAT AAT CGG AAT TAT CGT Met Lys Lys Tyr Leu Leu Gly Ile Gly Leu Ile Leu Ala Leu Ile Ala> 90 80 60 50 TGT AAG CAA AAT GTT AGC AGC CTT GAC GAG AAA-AAC AGC GTT TCA GTA ACA TTC GTT TTA CAA TCG TCG GAA CTG CTC TTT TTG TCG CAA AGT CAT Cys Lys Gln Asn Val Ser Ser Leu Asp Glu Lys Asn Ser Val Ser Val>

140 130 120 110 100

GAT TTG CCT GGT GAA ATG AAA GTT CTT GTA AGC AAA GAA AAA AAC AAA CTA AAC GGA CCA CTT TAC TTT CAA GAA CAT TCG TTT CTT TTT TTG TTT Asp Leu Pro Gly Glu Met Lys Val Leu Val Ser Lys Glu Lys Asn Lys>

180 170 160 150

GAC GGC AAG TAC GAT CTA ATT GCA ACA GTA GAC AAG CTT GAG CTT AAA CTG CCG TTC ATG CTA GAT TAA CGT TGT CAT CTG TTC GAA CTC GAA TTT Asp Gly Lys Tyr Asp Leu Ile Ala Thr Val Asp Lys Leu Glu Leu Lys>

230 220 210 200

GGA ACT TCT GAT AAA AAC AAT GGA TCT GGA GTA CTT GAA GGC GTA AAA CCT TGA AGA CTA TTT TTG TTA CCT AGA CCT CAT GAA CTT CCG CAT TTT Gly Thr Ser Asp Lys Asn Asn Gly Ser Gly Val Leu Glu Gly Val Lys>

> 270 280 260 250

GCT GAC AAA AGT AAA GTA AAA TTA ACA ATT TCT GAC GAT CTA GGT CAA CGA CTG TTT TCA TTT CAT TTT AAT TGT TAA AGA CTG CTA GAT CCA GTT Ala Asp Lys Ser Lys Val Lys Leu Thr Ile Ser Asp Asp Leu Gly Gln>

330 320 310 300 290

ACC ACA CTT GAA GTT TTC AAA GAA GAT GGC AAA ACA CTA GTA TCA AAA TGG TGT GAA CTT CAA AAG TTT CTT CTA CCG TTT TGT GAT CAT AGT TTT Thr Thr Leu Glu Val Phe Lys Glu Asp Gly Lys Thr Leu Val Ser Lys>

360 370 350 340

AAA GTA ACT TCC AAA GAC AAG TCA TCA ACA GAA GAA AAA TTC AAT GAA TTT CAT TGA AGG TTT CTG TTC AGT AGT TGT CTT TTT AAG TTA CTT Lys Val Thr Ser Lys Asp Lys Ser Ser Thr Glu Glu Lys Phe Asn Glu>

FIGURE 30 (1 of 4)

## B-31 OAP A/ B-31 OSP B / B-31 OSPC FUSION

חסכ		400	)		4:	10			420			43	Ģ.
390	•			•		•		•	•		•		•
AAA GGT GAA	GTA 1	rct (	SAA A	AAA A	ATA A	ATA	ACA .	AGA (	GCA	GAC	GGA	ACC	AGA
TTT CCA CTT	CAT	AGA C	TT :	TTT 3	TAT	TAT	TGT	TCT (	CGT	CTG	CCT	TGG	TCT
Lys Gly Glu	Val 9	Ser (	Glu I	Lvs	Ile :	Ile '	Thr	Arg :	Ala	Asp	Gly	Thr	Arg>
Lys diy did	102												_
440		4	450			46	0		4	70			480
•	,	•	•		•		•	•		•		•	•
CTT GAA TAC	ACA (	GGA A	ATT A	AAA A	AGC (	GAT (	GGA .	TCT (	GGA	AAA	GCT	AAA	GAG
CAA CTT ATG	TGT (	CCT :	TAA '	TTT :	TCG (	CTA	CCT	AGA-	CCT	TTT	CGA	TTT	CIC
Leu Glu Tyr	Thr	Gly :	Ile :	Lys :	Ser .	Asp	Gly	Ser	Gly	Lys	Ala	Lys	Glu>
		-											
4	90		5	00			510			52	20	•	
•	•	•	•	•		•	•		•		•	•	
GTT TTA AAA	GGC '	TAT	GTT	CTT	GAA	GGA	ACT	CTA	ACT	GCT	GAA	AAA	ACA
CAA AAT TTT	CCG .	ATA (	CAA	GAA	CTT	CCT	TGA	GAT	TGA.	CGA	CTT	TTT	TGT
Val Leu Lys	Gly	Tyr	Val	Leu	Glu	Gly	Thr	Leu	Thr	Ala	Glu	Lys	Thr>
					_						670		
530	540		_	55	0		3	60			570		•
• •			~ .	<b>663</b>	· CT	CTT.	» CT	ጥጥአ	) CC	**	a a T	איני	TCA
ACA TTG GTG	GTT	AAA	GAA	GGA	AC1	CVV	AC 1	777	TCG	444	TT A	WII	2 CT
TGT AAC CAC	CAA	TTT	CIT	CCI	Thr	Val	The	1.011	Ser	INS	Acn	Tle	Sers
Thr Leu Val	vai	Lys	Gin	GIY	1111	741	••••	200		٠, ٥	•••••		
580	9	90			600			61	.0		•	520	
580	,	90		•	•		•		•	•		•	
AAA TCT GGG	GAA	• GTT	TCA	• GTT	• GAA	CTT	• TAA	GAC	• ACT	• GAC	AGT	• AGT	GCT
AAA TCT GGC	GAA	GTT CAA	AGT	CAA	GAA CTT	GAA	TTA	GAC CTG	• ACT TGA	CTG	AGT TCA	AGT TCA	CGA
AAA TCT GGC	GAA	GTT CAA	AGT	CAA	GAA CTT	GAA	TTA	GAC CTG	• ACT TGA	CTG	AGT TCA	AGT TCA	CGA
AAA TCT GGG	GAA	GTT CAA	AGT	CAA	GAA CTT Glu	GAA Leu	TTA	GAC CTG	ACT TGA Thr	CTG	AGT TCA	AGT TCA Ser	CGA Ala>
AAA TCT GGC	GAA	GTT CAA	AGT Ser	CAA	GAA CTT Glu	GAA	TTA	GAC CTG	• ACT TGA	CTG	AGT TCA	AGT TCA Ser	CGA
AAA TCT GGC TTT AGA CCC Lys Ser Gly	G GAA CTT Glu	GTT CAA Val	AGT Ser	CAA Val	GAA CTT Glu	GAA Leu 650	TTA Asn	GAC CTG Asp	ACT TGA Thr 660	Asp	AGT TCA Ser	AGT TCA Ser	CGA Ala>
AAA TCT GGC TTT AGA CCC Lys Ser Gly 630	G GAA CTT Glu	GTT CAA Val	AGT Ser 10 • GCA	CAA Val	GAA CTT Glu TGG	GAA Leu 650 AAT	TTA Asn GAC	GAC CTG Asp	ACT TGA Thr 660	ASP AGC	AGT TCA Ser	AGT TCA Ser 6	CGA Ala> 70 ACA
AAA TCT GGC TTT AGA CCC Lys Ser Gly  630 GCT ACT AA	G GAA CTT Glu	GTT CAA Val 64 ACT TGA	AGT Ser 10 • GCA CGT	CAA Val GCT CGA	GAA CTT Glu TGG ACC	GAA Leu 650 AAT TTA	TTA Asn GAC CTG	GAC CTG Asp • AGT TCA	ACT TGA Thr 660 ACT TGA	ASP AGC TCG	AGT TCA Ser ACT	AGT TCA Ser 6'	CGA Ala> 70 ACA TGT
AAA TCT GGC TTT AGA CCC Lys Ser Gly 630	G GAA CTT Glu	GTT CAA Val 64 ACT TGA	AGT Ser 10 • GCA CGT	CAA Val GCT CGA	GAA CTT Glu TGG ACC	GAA Leu 650 AAT TTA	TTA Asn GAC CTG	GAC CTG Asp • AGT TCA	ACT TGA Thr 660 ACT TGA	ASP AGC TCG	AGT TCA Ser ACT	AGT TCA Ser 6'	CGA Ala> 70 ACA TGT
AAA TCT GGC TTT AGA CCC Lys Ser Gly  630  GCT ACT AA CGA TGA TT Ala Thr Ly	G GAA CTT Glu A AAA T TTT S Lys	GTT CAA Val 64 ACT TGA	AGT Ser 10 GCA CGT Ala	CAA Val GCT CGA	GAA CTT Glu TGG ACC	GAA Leu 650 AAT TTA Asn	TTA Asn GAC CTG Asp	GAC CTG Asp • AGT TCA	ACT TGA Thr 660 ACT TGA	ASP AGC TCG	AGT TCA Ser ACT	AGT TCA Ser 6'	CGA Ala> 70 ACA TGT
AAA TCT GGC TTT AGA CCC Lys Ser Gly  630 GCT ACT AA	G GAA CTT Glu A AAA T TTT S Lys	GTT CAA Val 64 ACT TGA	AGT Ser 10 • GCA CGT	CAA Val GCT CGA	GAA CTT Glu TGG ACC	GAA Leu 650 AAT TTA Asn	TTA Asn GAC CTG	GAC CTG Asp • AGT TCA	ACT TGA Thr 660 ACT TGA	AGC TCG Ser	AGT TCA Ser ACT	AGT TCA Ser 6'	CGA Ala> 70 ACA TGT Thr>
AAA TCT GGC TTT AGA CCC Lys Ser Gly 630 GCT ACT AAA CGA TGA TT Ala Thr Ly 680	G GAA C CTT Glu A AAA T TTT S Lys	GTT CAA Val 60 ACT TGA Thr	AGT Ser 10 GCA CGT Ala 690	CAA Val GCT CGA Ala	GAA CTT Glu TGG ACC Trp	GAA Leu 650 AAT TTA ASD	GAC CTG ASP	GAC CTG Asp AGT TCA Ser	ACT TGA Thr 660 ACT TGA Thr	AGC TCG Ser	AGT TCA Ser ACT TGA Thr	AGT TCA Ser 6' TTA AAT Leu	CGA Ala> 70 ACA TGT Thr> 720 GAT
AAA TCT GGC TTT AGA CCC Lys Ser Gly 630 GCT ACT AA CGA TGA TT Ala Thr Ly 680 ATT AGT GC	G GAA C CTT Glu A AAA T TTT S Lys T GAC	GTT CAA Val 60 ACT TGA Thr	AGT Ser 10 GCA CGT Ala 690 AAA	CAA Val GCT CGA Ala	GAA CTT Glu TGG ACC Trp	GAA Leu 650 AAT TTA ASD 7	GAC CTG Asp	GAC CTG Asp AGT TCA Ser	ACT TGA Thr 660 ACT TGA Thr	AGC TCG Ser 710	AGT TCA Ser ACT TGA Thr	AGT TCA Ser 6' TTA AAT Leu ACA	CGA Ala> 70 ACA TGT Thr> 720 GAT CTA
AAA TCT GGC TTT AGA CCC Lys Ser Gly 630 GCT ACT AA CGA TGA TT Ala Thr Ly 680 ATT AGT GC	G GAA C CTT Glu A AAA T TTT S Lys T GAC	GTT CAA Val 60 ACT TGA Thr	AGT Ser 10 GCA CGT Ala 690 AAA	CAA Val GCT CGA Ala	GAA CTT Glu TGG ACC Trp	GAA Leu 650 AAT TTA ASD 7	GAC CTG Asp	GAC CTG Asp AGT TCA Ser	ACT TGA Thr 660 ACT TGA Thr	AGC TCG Ser 710	AGT TCA Ser ACT TGA Thr	AGT TCA Ser 6' TTA AAT Leu ACA	CGA Ala> 70 ACA TGT Thr> 720 GAT CTA
AAA TCT GGC TTT AGA CCC Lys Ser Gly 630 GCT ACT AAA CGA TGA TT Ala Thr Ly 680	G GAA C CTT Glu A AAA T TTT S Lys T GAC	GTT CAA Val 60 ACT TGA Thr	AGT Ser 10 GCA CGT Ala 690 AAA	CAA Val GCT CGA Ala	GAA CTT Glu TGG ACC Trp	GAA Leu 650 AAT TTA ASD 7	GAC CTG Asp	GAC CTG Asp AGT TCA Ser	ACT TGA Thr 660 ACT TGA Thr	AGC TCG Ser 710 TTC	AGT TCA Ser ACT TGA Thr	AGT TCA Ser 6' TTA AAT Leu ACA	CGA Ala> 70 ACA TGT Thr> 720 GAT CTA
AAA TCT GGC TTT AGA CCC Lys Ser Gly 630 GCT ACT AA CGA TGA TT Ala Thr Ly 680 ATT AGT GC	G GAA C CTT Glu A AAA T TTT S Lys T GAC	GTT CAA Val 60 ACT TGA Thr	AGT Ser 10 GCA CGT Ala 690 AAA	CAA Val GCT CGA Ala	GAA CTT Glu TGG ACC Trp	GAA Leu 650 AAT TTA ASD 7	GAC CTG Asp	GAC CTG Asp AGT TCA Ser TTG AAC	ACT TGA Thr 660 ACT TGA Thr	AGC TCG Ser 710 TTC	AGT TCA Ser ACT TGA THA	AGT TCA Ser 6' TTA AAT Leu ACA	CGA Ala> 70 ACA TGT Thr> 720 GAT CTA
AAA TCT GGC TTT AGA CCC Lys Ser Gly  630  GCT ACT AAA CGA TGA TT Ala Thr Ly  680  ATT AGT GC TAA TCA CG Ile Ser Al	G GAA CTT Glu A AAA I TTT S Lys T GAC A CTG A ASP	GTT CAA Val 60 ACT TGA Thr AGC TCG Ser	AGT Ser 10 GCA CGT Ala 690 AAA TTT Lys	GCT CGA Ala AAA TTT Lys	GAA CTT Glu TGG ACC Trp	GAA Leu 650 AAT TTA ASD 7 AAA TTI Lys	GAC CTG ASP 00 GAT CTA S ASP	GAC CTG Asp AGT TCA Ser TTG AAC	ACT TGA Thr 660 ACT TGA Thr CAC	AGC TCG Ser 710 TTC AAC	AGT TCA Ser ACT TGA Thr TAA AAT Leu 160	AGT TCA Ser 6 TTA AAT Leu ACA TGT	CGA Ala> 70 ACA TGT Thr> 720 GAT CTA Asp>
AAA TCT GGC TTT AGA CCC Lys Ser Gly  630  GCT ACT AAA CGA TGA TT Ala Thr Ly  680  ATT AGT GC TAA TCA CG Ile Ser Al	G GAA CTT Glu A AAA TTTT S Lys T GAC A CTG A ASP	GTT CAA Val 60 ACT TGA Thr AGC TCG Ser	AGT Ser 10 GCA CGT Ala 690 AAA TTT Lys	GCT CGA Ala AAA TTT Lys	GAA CTT Glu TGG ACC TIP	GAA Leu 650 AAT TTA ASN 7 AAA TTT Lys	GAC CTG ASP 00 GAT CTA SASP	GAC CTG Asp AGT TCA Ser TTG AAC Leu	ACT TGA Thr 660 ACT TGA Thr CAO Val	AGC TCG Ser 710 TTCC AAC	AGT TCA Ser ACT TGA THA THA TABLE TA	AGT TCA Ser 6 TTA AAT Leu ACA TGT	CGA Ala> 70 ACA TGT Thr> 720 GAT CTA ASP>
AAA TCT GGC TTT AGA CCC Lys Ser Gly  630  GCT ACT AAA CGA TGA TT Ala Thr Ly  680  ATT AGT GC TAA TCA CG Ile Ser Al	G GAA CTT Glu A AAA TTTT S Lys T GAC A CTG A ASP	GTT CAA Val 60 ACT TGA Thr AGC TCG Ser	GCA GCA TITT Lys	GCT CGA Ala AAA TTT Lys 740	GAA CTT Glu TGG ACC Trp ACT TGA	GAA Leu 650 AAT TTA ASn 7 AAA TTT Lys	GAC CTG ASP 00 GAT CTA GAT CTA GAT GAT GAT GAT GAT GAT GAT GAT	GAC CTG Asp  AGT TCA Ser TTG AAC Leu A GCT	ACT TGA Thr 660 ACT TGA Thr CAO Val	AGC TCG Ser 710 TTCC AACC Phe	AGT TCA Ser ACT TGA Thr TAA AAT AAAT AAAT AAAT AAAT AAAT A	AGT TCA Ser 6 TTA AAT Leu ACA TGI Thi	CGA Ala> 70 ACA TGT Thr> 720 GAT CTA ASP> A GAA CTT

FIGURE 30 (2 of 4)

### B-31 OSP A/ B-31 OSP B / B-31 OSP C FUSION

770			780			79	0		8	00			810		
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CC2	TCA	GCA	AGT	GAA	ATT	AAA	TAA	CTT	TCA	GAG	CTT	AAA	AAC	GCT	TTA
CCT	) CT	CCT	TCA	CAL	TAA	TTT	TTA	GAA	AGT	CTC	GAA	TIT	TTG	CGA	33T
COL	VOI	*15	202	Glu	Tla	INS	Asn	Leu	Ser	Glu	Leu	LVS	255	212	Leu>
GIA	Sei	VIC	Ser	Giu	-16	טייט	••••					-3-		****	2607
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82	20			330		_	040			٠.	*		•	-	
•	•	•						C1.	-		101	-			
AAA	ATG	GCT	TAA	AAT	TCA	GGG	AAA	GAI	666	AAI	ACA	101	GCA	AAT	TCT
TTT	TAC	CGA	TTA	TTA	AGT	CCC	TTT	CTA	CCC	TTA	TGT	AGA	CGT	ATT	AGA
Lys	Met	Ala	Asn	Asn	Ser	Gly	Lys	Asp	GIA.	ASD.	.Thr	Ser	Ala	Asn	Ser>
	870			88	30		8	90			900			93	10
•	•		•		•	•		•		• '	•		• •	••	• .
GCT	GAT	GAG	TCT	GTT	AAA	GGG	CCT	TAA	CTT	ACA	GAA	ATA	λGT	አጹኣ	AAA
CGA	CTA	CTC	AGA	CAA	TTT	CCC	GGA	TTA	GAA	TGT	CTT	TAT	TCA	TTT	LIL
Ala	Asp	Glu	Ser	Val	Lys	Gly	Pro	Asn	Leu	Thr	Glu	Ile	Ser	Lys	Lys>
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	, ,	920			930			94	0		9	950			960.
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TTA	ACG	GAT	тст	TAA	GCG	GTT	TTA	CTT	GCT	GTG	AAA	GAG	GTT	GAA	GCG
AAT	TGC	CTA	AGA	TTA	CGC	CAA	AAT	GAA	CGA	CAC	TTT	CTC	CAA	CTT	CGC
Tle	Thr	'Asp	Ser	Asn	Ala	Val	Leu	Leu	Ala	Val	Lys	Glu	Val	Glu	Ala>
	••••										-				
		9	70		9	980			990			10	00		
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<b>ጥ</b> ፓር	СТС	ጥርኔ	тст	ATA	GAT	GAA	TTA	GCT	GCT	AAA	GCT	ATT	GGT	AAA	<del>አ</del> ዱአ
220	CAC	) ACT	707	TAT	CTA	CTT	TAA	CGA	CGA	TTT	CGA	AAT	CCA	TTT	TTT
LAU	IAU	Ser	Ser	Tle	Asp	Glu	Ile	Ala	Ala	Lys	Ala	Ile	Glv	Lvs	Lys>
Leu	Deu		501							•				•	-•
1010			1020			10	30		1	040			1050		
1010		•	1020		•		•	•		•		•	•		•
N TO N	CNC	CAN	) እስጥ	አኔጥ	CCT	ттс	GAT	ACC	GAA	TAT	ААТ	CAC	AAT	GGA	TCA
WIN	CAC	CAM	, ww.,	TTA	CCN	220	CTA	TGG	بلمل	מדמ	TTA	CTG	TTA	CCT	AGT
TAI	610	011	. 114	1114	Clar	tou	Acn	Thr	Glu	Tyr	) len	Hic	Acn	Glv	Ser>
116	: 112	GII	I ASI	ASII	Gry	Deu	, vab		014	•3•	*****		****	O-,	
			•	070			1080			10	90		1	100	
10	060		. 1	.070		•	1000		•		•		•		
				-			እጥጸ	ጥር እ	» CC	· ~~	አጥአ	* * *	Chh	2 2 2	TTA
TTC	; TT/	A GCC	, GGP	, CG1	IAI	CCT	. wyw , win	NOT	י אכנ	CAT	, <i>wyw</i>	. 4444	. CTT	4444.	TAA
AAC	AA:	r CGG	CCI	GCA	ATA		TAL	WGT	. w.	GAI	TAL		Cla	111	LAUS
Lev	ı Lei	ı Ala	3 G17	Arg	тУI	. ATS	, TT6	. ser	1111	. Let	· TTG	- Lys	GIU	- Lys	Leu>
		_					,	120			1140			11	.50
	111	0		11	.20		. ,	.130		•	1140			11	
•		•	•												
GA'	T GG	A TT	g aa	A AA?	GA	A GG/	4 TT7	AAC	, GA	1 AA/	. All	GAI	GC0	, 661	A÷G
CT.	A CC	AA T	CTT	r TT	\ CT	r CC.	r AA7	. 110	. CF	1 1-1".	TAA	CTA	CGC	. CGA	TIC
As	p G1	y Le	u Ly	s Ası	ı Glı	1 G1;	y Lei	ı Ly:	5 GI	ı Ly:	2 116	ASĮ	) Ala	. Ala	Lys>

FIGURE 30 (3 of 4)

#### B-31 OSP A / B-31 OSP B/ B-31 OSP C FUSION

1190 1200 1180 1160 AAA TGT TCT GAA ACA TTT ACT AAT AAA TTA AAA GCA AAA CAC ACA GAT TTT ACA AGA CTT TGT AAA TGA TTA TTT AAT TTT CGT TTT GTG TGT CTA Lys Cys Ser Glu Thr Phe Thr Asn Lys Leu Lys Ala Lys Ris Thr Asp> 1240 1230 1220 1210 CTT GGT AAA GAA GGT GTT ACT GAT GCT GAT GCA AAA GAA GCC ATT TTA GAA CCA TTT CTT CCA CAA TGA CTA CGA CTA CGT TTT CTT CGG TAA AAT Leu Gly Lys Glu Gly Val Thr Asp Ala Asp Ala Lys Glu Ala Ile Leu> 1280 1290 1270 1260 AAA ACA AAT GGT ACT AAA ACT AAA GGT GCT GAA GAA CTT GGA AAA TTA TTT TGT TTA CCA TGA TTT TGA TTT CCA CGA CTT CTT GAA CCT TTT AAT Lys Thr Asn Gly Thr Lys Thr Lys Gly Ala Glu Glu Leu Gly Lys Leu> 1320 1330 1310 TTT GAA TCA GTA GAG GTC TTG TCA AAA GCA GCT AAA GAG ATG CTT GCT AAA CTT AGT CAT CTC CAG AAC AGT TTT CGT CGA TTT CTC TAC GAA CGA Phe Glu Ser Val Glu Val Leu Ser Lys Ala Ala Lys Glu Met Leu Ala> 1380 1370 1360 1350 • AAT TCA GTT AAA GAG CTT ACA AGC CCT GTT GTG GCA GAA AGT CCA AAA TTA AGT CAA TTT CTC GAA TGT TCG GGA CAA CAC CGT CTT TCA GGT TTT Asn Ser Val Lys Glu Leu Thr Ser Pro Val Val Ala Glu Ser Pro Lys>

1400

AAA CCT TAA TTT GGA ATT Lys Pro ***>

#### B-31 OSP C/ B-31 OSP A/ B-31 OSP B FUSION

		,	10			20			30			4	0		
	•	•	•	•		•		•	•		•		•	•	
ATG	AAA	AAG	AAT	ACA	TTA	AGT	GCG	ATA	ATT	ATG	ACT	TTA	TTT	TTA	TTT
TAC	TTT	TTC	TTA	TGT	AAT	TCA	CGC	TAT	TAA	TAC	TGA	AAT	AAA	TAA	<b>KAA</b>
Met	Lys	Lys	Asn	Thr	Leu	Ser	Ala	Ile	Leu	Met	Thr	Leu	Phe	Leu	Phe>
	-	-							•						
50			60			7	0			80		_	90		
•		•	•		•		•	~ ~			101		•		
ATA	TCT	TGT	AAT	TAA	TCA	GGG	AAA	CTA	CCC	WAT.	ACA	101	CCT	AAT	TCT
TAT	AGA	ACA	ATT	TTA	AGT	CIV	tre	CIN	GIV	Jen-	TGT Thr	707	γla	1114	AGA Com
He	Ser	Cys	ASn	ASII	Ser	GIY	Lys	vab	O13	11311	••••		n1a		2617
10	۱۸		1	10			120			13	30		1	140	••
10	•	•		•		•	•		•		•	•		•	
GCT	GAT	GAG	TCT	GTT	AAA	GGG	CCT	TAA	CTT	ACA	GAA	ATA	AGT	AAA	AAA
CGA	СТА	CTC	AGA	CAA	TTT	CČC	GGA	ATT	GAA	TGT	CTT	TAT	TCA	TIT	TTT
Ala	Asp	Glu	Ser	Val	Lys	Gly	Pro	Asn	Leu	Thr	Glu	Ile	Ser	<u>:</u> ys	Lys>
				•				70			100				
_	150		_	16	50	•		170		•	180		•	19	•
7 T-T	»CC	C N T	TCT.	דממ	GCG	CTT	TTA	CTT	GCT	GTG	AAA	GAG	GTT	GAA	GCG
TAA	TCC	CTA	)C)	ፈጥጉ ሊጉጉ	CGC	CAA	AAT	GAA	CGA	CAC	TTT	CTC	CAA	CTT	CGC
Tle	Thr	ASD	Ser	Asn	Ala	Val	Leu	Leu	Ala	Val	Lys	Glu	Val	Glu	Ala>
		200			210			2:	20			230			240
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TTG	CTG	TCA	TCT	ATA	GAT	GAA	TTA	GCT	GCT	AAA	GCT	ATT	GGT	ኢኢኒ ኤኤኤ	33.3 33.3
AAC	GAC	AGT	AGA	TAT	CTA	CIT	TAA	אטט	A I a	111	CGA	TIAA	CLA	111	Lys>
Leu	Leu	Ser	Ser	116	ASD	GIU	116	VIG	VIG	Lys	VIG	116	GIY	Lys	Lys
		2	50			260			270			2	80		
	•	-	•	÷		•		•	•		•		•	•	
ATA	CAC	CAA	TAA	TAA	GGT	TTG	GAT	ACC	GAA	TAT	AAT	CAC	TAA	GGA	TCA
TAT	GTG	GTT	ATT 7	ATT .	CCA	AAC	CTA	TGG	CTT	ATA	ATT	GTG	TTA	CCT	AGT
Ile	His	Glr	ı Asn	Asn	Gly	Leu	q2A	Thr	Glu	Tyr	Asn	His	Asn	Gly	Ser>
			200			7	10			320			330		
290			300	,	•	,	•	•		*		•	•		•
TTG	TT.	GCC	G GGZ	CGT	TAT	GCA	ATA	TCA	ACC	CTA	ATA .	. AAA	CAA	AAA	TTA
AAC	: AA1	CG(	c cc1	r GCA	ATA	CGI	TAT 1	, YCI	TGG	GAT	TAT :	, LLI	GI1	, TII	TAA
Leu	ı Lei	ı Al	a Gly	Arg	Tyr	Ala	1le	Ser	Thr	Leu	Ile	Lys	Glr	Lys	reas
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#### B-31 OSP C/ B-31 OSP A/ B-31 OSP B FUSION

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		TĊT													
		AGA													
Lys	Cys	Ser	Glu	Thr	Phe	Thr	Asn	Lys	Leu	Lys	Ala	Lys	His	Thr	Asp>
	4	140			450			46	50	_	4	170			480
•				•	•	·			•					•	•
		AAA													
		TTT													
Leu	Gly	Lys	GIU	GIA	Vai	Inr	ASD	Ala	ASP	AIG	Lys	GIU	AIA	TIE	Leu>
		49	0		9	00			510			52	20		
	•		•	•		•	•	•	•		•		•	٠. •	
AAA	ACA	AAT	GGT	ACT	AAA	ACT	AAA	GGT	GCT	GAA	GAA	CTT	GGA	AAA	TTA
TTT	TGT	TTA	CCA	TGA	TTT	TGA	TTT	CCA	CGA	CTT	CTT	GAA	CCT	TTT	AAT
Ĺys	Thr	Asn	Gly	Thr	Lys	Thr	Lys	Gly	Ala	Glu	Gļu	Leu	Gly	Lys	Leu>
530			540			55	50		9	60			570		
•		•	•		•		•	•		•		•	1 *		•
TTT	GAA	TCA	GTA	GAG	GTC	TTG	TCA	AAA	GCA	GCT	AAA	GAG	'ATG	CTT	GCT
AAA	CTT	AGT	CAT	CTC	CAG	AAC	AGT	TTT	CGT	CGA	TTT	CTC	TAC	GAA	CGA
Phe	Glu	Ser	Val	Glu	Val	Leu	Ser	Lys	Ala	Ala	Lys	Glu	Met	Leu	Ala>
58	30		9	590			600			61	10		6	20	*
58	30	•	:	590		•	600 •		•	61	LO •	•	•	20	
ТАА	TCA	• GTT	AAA	• GAG			* AGC			GTG	GCA		AGT	• CCA	
TAA ATT	TCA AGT	CAA	AAA TTT	GAG CTC	GAA	TGT	AGC TCG	GGA	CAA	GTG CAC,	GCA CGT	CTT	AGT TCA	CCA GGT	TTT
TAA ATT	TCA AGT	CAA	AAA TTT	GAG CTC	GAA	TGT	AGC TCG	GGA	CAA	GTG CAC,	GCA CGT	CTT	AGT TCA	CCA GGT	
TAA ATT	TCA AGT	CAA	AAA TTT	GAG CTC Glu	GAA	TGT	AGC TCG Ser	GGA	CAA	GTG CAC,	GCA CGT	CTT	AGT TCA	CCA GGT	TTT Lys>
AAT TTA Asn	TCA AGT Ser	CAA Val	AAA TTT Lys	GAG CTC Glu	GAA Leu 40	TGT Thr	AGC TCG Ser	GGA Pro 650	CAA Val	GTG CAC, Val	GCA CGT Ala	CTT Glu	AGT TCA Ser	CCA GGT Pro	TTT Lys>
TAA ATT Asn	TCA AGT Ser 630	CAA Val	AAA TTT Lys •	GAG CTC Glu 64	GAA Leu 40 • GTT	TGT Thr	AGC Ser	GGA Pro 650 • CTT	CAA Val GAC	GTG CAC, Val	GCA CGT Ala 660	CTT Glu AAC	AGT TCA Ser	CCA GGT Pro 67	TTT Lys> 10 TCA
AAT TTA Asn • • • • • • • • • • • • • • • • • • •	TCA AGT Ser 630 CCT GGA	CAA Val AAG TTC	AAA TTT Lys CAA GTT	GAG CTC Glu 64 AAT TTA	GAA Leu 40 • GTT CAA	TGT Thr AGC TCG	AGC TCG Ser AGC TCG	GGA Pro 650 CTT GAA	CAA Val GAC CTG	GTG CAC, Val	GCA CGT Ala 660 AAA TTT	CTT Glu AAC TTG	AGT TCA Ser AGC TCG	CCA GGT Pro 67 GTT CAA	TTT Lys> 10 TCA AGT
AAT TTA Asn • • • • • • • • • • • • • • • • • • •	TCA AGT Ser 630 CCT GGA	CAA Val AAG TTC	AAA TTT Lys CAA GTT	GAG CTC Glu 64 AAT TTA	GAA Leu 40 • GTT CAA	TGT Thr AGC TCG	AGC TCG Ser AGC TCG	GGA Pro 650 CTT GAA	CAA Val GAC CTG	GTG CAC, Val	GCA CGT Ala 660 AAA TTT	CTT Glu AAC TTG	AGT TCA Ser AGC TCG	CCA GGT Pro 67 GTT CAA	TTT Lys> 10 TCA
AAT TTA Asn • • • • • • • • • • • • • • • • • • •	TCA AGT Ser 630 CCT GGA Pro	CAA Val AAG TTC	AAA TTT Lys CAA GTT	GAG CTC Glu 64 AAT TTA	GAA Leu 40 • GTT CAA	TGT Thr AGC TCG	AGC TCG Ser AGC TCG	GGA Pro 550 CTT GAA Leu	CAA Val GAC CTG	GTG CAC, Val	GCA CGT Ala 660 AAA TTT Lys	CTT Glu AAC TTG	AGT TCA Ser AGC TCG	CCA GGT Pro 67 GTT CAA	TTT Lys> 10 TCA AGT
AAT TTA Asn • • • • • • • • • • • • • • • • • • •	TCA AGT Ser 630 CCT GGA Pro	CAA Val AAG TTC Lys	AAA TTT Lys CAA GTT	GAG CTC Glu 64 AAT TTA	GAA Leu 40 GTT CAA Val	TGT Thr AGC TCG	AGC TCG Ser AGC TCG	GGA Pro 550 CTT GAA Leu	CAA Val GAC CTG Asp	GTG CAC, Val	GCA CGT Ala 660 AAA TTT Lys	CTT Glu AAC TTG Asn	AGT TCA Ser AGC TCG	CCA GGT Pro 67 GTT CAA	TTT Lys> 70 • TCA AGT Ser>
AAT TTA Asn AAA TTT Lys	TCA AGT Ser 630 CCT GGA Pro	CAA Val AAG TTC Lys	AAA TTT Lys CAA GTT Gln	GAG CTC Glu 64 AAT TTA Asn	GAA Leu 40 GTT CAA Val	TGT Thr AGC TCG Ser	AGC TCG Ser AGC TCG Ser	GGA Pro 650 CTT GAA Leu	CAA Val GAC CTG Asp	GTG CAC, Val GAG CTC Glu	GCA CGT Ala 660 AAA TTT Lys	AAC TTG Asn	AGT TCA Ser AGC TCG Ser	CCA GGT Pro 67 GTT CAA Val	TTT Lys> 70 TCA AGT Ser>
AAT TTA Asn AAA TTT Lys GTA CAT	TCA AGT Ser 630 CCT GGA Pro	AAG TTC Lys 680	AAA TTT Lys CAA GTT Gln	GAG CTC Glu 64 AAT TTA Asn	GAA Leu 10 GTT CAA Val 690 GAA CTT	TGT Thr AGC TCG Ser ATG TAC	AGC TCG Ser AGC TCG Ser AAA TTT	GGA Pro 650 CTT GAA Leu 7 GTT CAA	GAC CTG Asp	GTG CAC, Val GAG CTC Glu GTA CAT	GCA CGT Ala 660 AAA TTT Lys	AAC TTG ASN	AGT TCA Ser AGC TCG Ser GAA CTT	CCA GGT Pro 6T GTT CAA Val	TTT Lys> 70 TCA AGT Ser> 720 AAC
AAT TTA Asn AAA TTT Lys GTA CAT	TCA AGT Ser 630 CCT GGA Pro	AAG TTC Lys 680	AAA TTT Lys CAA GTT Gln	GAG CTC Glu 64 AAT TTA Asn	GAA Leu 10 GTT CAA Val 690 GAA CTT	TGT Thr AGC TCG Ser ATG TAC	AGC TCG Ser AGC TCG Ser AAA TTT	GGA Pro 650 CTT GAA Leu 7 GTT CAA	GAC CTG Asp	GTG CAC, Val GAG CTC Glu GTA CAT	GCA CGT Ala 660 AAA TTT Lys	AAC TTG ASN	AGT TCA Ser AGC TCG Ser GAA CTT	CCA GGT Pro 6T GTT CAA Val	TTT Lys> 70 TCA AGT Ser> 720
AAT TTA Asn AAA TTT Lys GTA CAT	TCA AGT Ser 630 CCT GGA Pro	AAG TTC Lys 680 TTG AAC	AAA TTT Lys CAA GTT Gln	GAG CTC Glu 64 AAT TTA Asn	GAA Leu 40 GTT CAA Val 690 GAA CTT Glu	TGT Thr AGC TCG Ser ATG TAC	AGC TCG Ser AGC TCG Ser AAA TTT	GGA Pro 650 CTT GAA Leu 7 GTT CAA	GAC CTG Asp	GTG CAC, Val GAG CTC Glu GTA CAT Val	GCA CGT Ala 660 AAA TTT Lys	AAC TTG Asn 710 AAA TTT Lys	AGT TCA Ser AGC TCG Ser GAA CTT	CCA GGT Pro 6T GTT CAA Val	TTT Lys> 70 TCA AGT Ser> 720 AAC
AAT TTA ASD AAA TTT Lys GTA CAT Val	TCA AGT Ser 630 CCT GGA Pro	AAG TTC Lys 680 TTG AAC Leu	AAA TTT Lys CAA GTT Gln CCT GGA Pro	GAG CTC Glu 64 AAT TTA ASD GGT CCA Gly	GAA Leu 10 GTT CAA Val 690 GAA CTT Glu	TGT Thr AGC TCG Ser ATG TAC Met	AGC TCG Ser AGC TCG Ser AAA TTT Lys	GGA Pro 650 CTT GAA Leu 7 GTT CAA Val	GAC CTG Asp OCTT GAA Leu	GTG CAC, Val GAG CTC Glu GTA CAT Val	GCA CGT Ala 660 AAA TTT Lys AGC TCG Ser	AAC TTG Asn 710 AAA TTT Lys	AGT TCA Ser AGC TCG Ser GAA CTT Glu	CCA GGT Pro 67 GTT CAA Val AAA TTT Lys	TTT Lys> 70 TCA AGT Ser> 720 AAC TTG Asn>
AAT TTA ASD AAA TTT Lys GTA CAT Val	TCA AGT Ser 630 CCT GGA Pro	AAG TTC Lys 680 TTG AAC Leu 7	AAA TTT Lys CAA GTT Gln CCT GGA Pro	GAG CTC Glu 64 AAT TTA ASN GGT CCA Gly	GAA Leu  10 GTT CAA Val 690 GAA CTT Glu	TGT Thr AGC TCG Ser ATG TAC Met	AGC TCG Ser AGC TCG Ser Lys	GGA Pro 650 CTT GAA Leu 7 GTT CAA Val	GAC CTG Asp OCTT GAA Leu 750	GTG CAC, Val GAG CTC Glu GTA CAT Val	GCA CGT Ala 660 AAA TTT Lys AGC TCG Ser	AAC TTG Asn 710 AAA TTT Lys 7	AGT TCA Ser AGC TCG Ser GAA CTT Glu	CCA GGT Pro 67 GTT CAA Val AAA TTT Lys	TTT Lys> 70 TCA AGT Ser> 720 AAC TTG ASN>
AAT TTA ASD  AAA TTT Lys  GTA CAT Val	TCA AGT Ser 630 CCT GGA Pro	AAG TTC Lys 680 TTG AAC Leu 7	AAA TTT Lys CAA GTT Gln CCT GGA Pro	GAG CTC Glu  64 AAT TTA ASN  CCA Gly  TAC	GAA Leu  10 GTT CAA Val 690 GAA CTT Glu GAT CTA	AGC TCG Ser ATG TAC Met CTA GAT	AGC TCG Ser AGC TCG Ser AAA TTT Lys	GGA Pro 650 CTT GAA Leu 7 GTT CAA Val	GAC CTG ASP 00 CTT GAA Leu 750 ACA	GTG CAC, Val GAG CTC Glu GTA CAT Val	GCA CGT Ala 660 AAA TTT Lys AGC TCG Ser	AAC TTG ASn TTT Lys 7	AGT TCA Ser AGC TCG Ser GAA CTT Glu	CCA GGT Pro 67 GTT CAA Val AAA TTT Lys	TTT Lys> 70 TCA AGT Ser> 720 AAC TTG Asn>
AAT TTA ASD  AAA TTT Lys  GTA CAT Val	TCA AGT Ser 630 CCT GGA Pro	AAG TTC Lys 680 TTG AAC Leu 7	AAA TTT Lys CAA GTT Gln CCT GGA Pro	GAG CTC Glu  64 AAT TTA ASN  CCA Gly  TAC	GAA Leu  10 GTT CAA Val 690 GAA CTT Glu GAT CTA	AGC TCG Ser ATG TAC Met CTA GAT	AGC TCG Ser AGC TCG Ser AAA TTT Lys	GGA Pro 650 CTT GAA Leu 7 GTT CAA Val	GAC CTG ASP 00 CTT GAA Leu 750 ACA	GTG CAC, Val GAG CTC Glu GTA CAT Val	GCA CGT Ala 660 AAA TTT Lys AGC TCG Ser	AAC TTG ASn TTT Lys 7	AGT TCA Ser AGC TCG Ser GAA CTT Glu	CCA GGT Pro 67 GTT CAA Val AAA TTT Lys	TTT Lys> 70 TCA AGT Ser> 720 AAC TTG ASN>

FIGURE 31 (2 of 4)

 $\langle \dot{\gamma} \rangle$ 

65/133

# B-31 OSP C/ B-31 OSP A/ B-31 OSP B FUSION

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			203	$\sim$	TATA	TTC	TTA	CCL	AGA	cc1	CAI		CII		CVI
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TTT	CGA	CTG	TTT	TCA	TTT	CAT	TIT	AAT	TGT	TAA	AGA	CIG	CIA	CAL	CLA
Lys	Ala	Asp	Lys	Ser	Lys	Val	Lys	Leu	Thr	11.6	Ser	wab	wsb	Let.	Gly>
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		TOT.	CAA	CTT	CAA	AAG	TTT	CTT	CTA	CCG	111,	161	GAI	CA.	MG I
GTT	166	101	Lan	Glu	Val	Phe	Lvs	Glu	Asp	Gly	Lys	Thr	Leu	Val	Ser>
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TC	r ga	A CT	T AT	G TG	I CC	T TA	A TT	TTO	G CI	A CC	1 WO	. CC.	t Lv:	s Ala	Lys>
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FIGURE 31 (3 of 4)

B-31 OSP C/ B-31 OSP A / B-31 OSP B FUSION

1170 1180 1190 1200 TCA AAA TCT GGG GAA GTT TCA GTT GAA CTT AAT GAC ACT GAC AGT AGT AGT TTT AGA CCC CTT CAA AGT CAA CTT GAA TTA CTG TGA CTG TCA TCA Ser Lys Ser Gly Glu Val Ser Val Glu Leu Asn Asp Thr Asp Ser Ser> 1210 1220 1230 1240 GCT GCT ACT AAA AAA ACT GCA GCT TGG AAT GAC AGT ACT AGC ACT TTA CGA CGA TGA TTT TTT TGA CGT CGA ACC TTA CTG TCA TGA TCG TGA AAT Ala Ala Thr Lys Lys Thr Ala Ala Trp Asn Asp Ser Thr Ser Thr Leus 1250 1260 1270 1280 ACA ATT AGT GCT GAC AGC AAA AAA ACT AAA GAT TTG GTG TTC TTA ACA TGT TAA TCA CGA CTG TCG TTT TTT TGA TTT CTA AAC CAC AAG AAT TGT Thr Ile Ser Ala Asp Ser Lys Lys Thr Lys Asp Leu Val Phe Leu Thr> 1300 1310 1320 1230 1340 GAT GGT ACA ATT ACA GTA CAA CAA TAC AAC ACA GCT GGA ACC AGC CTA CTA CCA TGT TAA TGT CAT GTT GTT ATG TTG TGT CGA CCT TGG TCG GAT Asp Gly Thr Ile Thr Val Gln Gln Tyr Asn Thr Ala Gly Thr Ser Leu> 1360 1370 1380 1390 GAA GGA TCA GCA AGT GAA ATT AAA AAT CTT TCA GAG CTT AAA AAC GCT

CTT CCT AGT CGT TCA CTT TAA TTT TTA GAA AGT CTC GAA TTT TTG CGA Glu Gly Ser Ala Ser Glu Ile Lys Asn Leu Ser Glu Leu Lys Asn Ala>

1400

TTA AAA TAA AAT TTT ATT Leu Lys ***>

#### **FUSION SEQUENCE**

19.15

B-31 OSP A/ B-31 P-93 (1168-2100) Sequence Range: 1 to 1720

> 10 ' 20 30 AAG CAA AAT GTT AGC AGC CTT GAC GAG AAA AAC AGC GTT TCA GTA GAT TTC GTT TTA CAA TCG TCG GAA CTG CTC TTT TTG TCG CAA AGT CAT CTA K Q N V S S L D E K N S V S · V D> 70 50 60 . 80_--. • TTG CCT GGT GAA ATG AAA GTT CTT GTA AGC AAA GAA AAA AAC AAA GAC AAC GGA CCA CTT TAC TTT CAA GAA CAT TCG TTT CTT TTT TTG TTT. CTG L P G E M K V L V S K E K N K D> 100 110 120 130 GGC AAG TAC GAT CTA ATT GCA ACA GTA GAC AAG CTT GAG CTT AAA GGA CCG TTC ATG CTA GAT TAA CGT TGT CAT CTG TTC GAA CTC GAA TTT CCT G K Y D L I A T V D K L E L K G> 170 160 · 150 180 ACT TCT GAT AAA AAC AAT GGA TCT GGA GTA CTT GAA GGC GTA AAA GCT TGA AGA CTA TTT TTG TTA CCT AGA CCT CAT GAA CTT CCG CAT TTT CGA T S D K N N G S G V L E G V K A> 220 200 210 230 240 GAC AAA AGT AAA GTA AAA TTA ACA ATT TCT GAC GAT CTA GGT CAA ACC CTG TTT TCA TTT CAT TTT AAT TGT TAA AGA CTG CTA GAT CCA GTT TGG D K S K V K L T I S D D L G Q T> 260 270 • ACA CTT GAA GTT TTC AAA GAA GAT GGC AAA ACA CTA GTA TCA AAA AAA TGT GAA CTT CAA AAG TTT CTT CTA CCG TTT TGT GAT CAT AGT TTT TTT T L E V F K E D G K T L V S K K> 290 300 310 . 320 330 • GTA ACT TCC AAA GAC AAG TCA TCA ACA GAA GAA AAA TTC AAT GAA AAA CAT TGA AGG TTT CTG TTC AGT AGT TGT CTT TTT AAG TTA CTT TTT V T S K D K S S T E E K F N E K> 340 350 360 370 380 GGT GAA GTA TCT GAA AAA ATA ATA ACA AGA GCA GAC GGA ACC AGA CTT CCA CTT CAT AGA CTT TTT TAT TAT TGT TCT CGT CTG CCT TGG TCT GAA G E V S E K I I T R A D G T R L>

> > FIGURE 32 (1 of 5)

B-31 OSP A/ B-31 P93

400 410 420 390 GAA TAC ACA GGA ATT AAA AGC GAT GGA TCT GGA AAA GCT AAA GAG GTT CTT ATG TGT CCT TAA TTT TCG CTA CCT AGA CCT TTT CGA TTT CTC CAA EYTGIKSDGSGKAKEV> 450 460 470 440 TTA AAA GGC TAT GTT CTT GAA GGA ACT CTA ACT GCT GAA AAA ACA ACA AAT TIT CCG ATA CAA GAA CTT CCT TGA GAT TGA CGA CTT TTT TGT TGT L K G Y V L E G T L T A E K T T> 490 500 510 520 • • • TTG GTG GTT AAA GAA GGA ACT GTT ACT TTA AGC AAA AAT ATT TCA AAA AAC CAC CAA TTT CTT CCT TGA CAA TGA AAT TCG TTT TTA TAA AGT TTT L V V K E G T V T L S K N I S K> 550 560 540 570 TCT GGG GAA GTT TCA GTT GAA CTT AAT GAC ACT GAC AGT AGT GCT GCT AGA CCC CTT CAA AGT CAA CTT GAA TTA CTG TGA CTG TCA TCA CGA CGA S G · E V S V E L N D T D S S A A> 600 580 590 610 ACT AAA AAA ACT GCA GCT TGG AAT TCA GGC ACT TCA ACT TTA ACA ATT TGA TIT TIT TGA CGT CGA ACC TTA AGT CCG TGA AGT TGA AAT TGT TAA T K K T A A W N S G T S T L T I>. 630 640 650 660 ACT GTA AAC AGT AAA AAA ACT AAA GAC CTT GTG TTT ACA AAA GAA AAC TGA CAT TTG TCA TTT TTT TGA TTT CTG GAA CAC AAA TGT TTT CTT TTG T V N S K K T K D L V F T K E N> 680 690 700 710 710 ACA ATT ACA GTA CAA CAA TAC GAC TCA AAT GGC ACC AAA TTA GAG GGG TGT TAA TGT CAT GTT ATG CTG AGT TTA CCG TGG TTT AAT CTC CCC T I T V Q Q Y D S N G T K L E G> 760 730 740 750 TCA GCA GTT GAA ATT ACA AAA CTT GAT GAA ATT AAA AAC GCT TTA AAA AGT CGT CAA CTT TAA TGT TTT GAA CTA CTT TAA TTT TTG CGA AAT TTT S A V E I T K L D E I K N A L K>

FIGURE 32 (2 of 5)

B-31 OSP A/ B-31 P-93

780 790 800 810 770 GGT CAC CCC ATG GAT GAA AAG CTT TTA AAA AGT AAA GAT GAT AAA GCA CCA GTG GGG TAC CTA CTT TTC GAA AAT TTT TCA TTT CTA CTA TTT CGT G H P M D E K L L K S K D D K A> 850 840 830 820 AGT AAA GAT GGT AAA GCC TTG GAT CTT GAT CGA GAA TTA AAT TCT AAA TCA TIT CTA CCA TIT CGG AAC CTA GAA CTA GCT CTT AAT TTA AGA TIT S K D G K A L D L D R R E E L N S K 880 890 900 GCT TCT AGC AAA GAA AAA AGT AAA GCC AAG GAA GAA ATA ACC AAG CGA AGA TCG TTT CTT TTT TCA TTT CGG TTC CTT CTT TAT TGG TTC A S S K E K S K A K E E E I T K> 940 950 930 920 • • • GGT AAG TCA CAG AAA AGC TTA GGC GAT TTG AAT AAT GAT GAA AAT CTT CCA TTC AGT GTC TTT TCG AAT CCG CTA AAC TTA TTA CTA CTT. TTA GAA G K S Q K S L G D L N N D E N L> 980 990 970 • ATG ATG CCA GAA GAT CAA AAA TTA CCT GAG GTT AAA AAA TTA GAT AGC TAC TAC GGT CTT CTA GTT TTT AAT GGA CTC CAA TTT TTT AAT CTA TCG M M P E D Q K L P E V K K L D S> 1020 1030 1040 1040 1050 AAA AAA GAA TTT AAA CCT GTT TCT GAG GTT GAG AAA TTA GAT AAG ATT TTT TTT CTT AAA TTT GGA CAA AGA CTC CAA CTC TTT AAT CTA TTC TAA K K E F K P V S E V E K L D K I> 1090 1070 1080 1100 TTC AAG TCT AAT AAC AAT GTT GGA GAA TTA TCA CCG TTA GAT AAA TCT AAG TTC AGA TTA TTG TTA CAA CCT CTT AAT AGT GGC AAT CTA TTT AGA F K S N N N V G E L S P L D K S> 1110 1120 1130 1140 TCT TAT AAA GAC ATT GAT TCA AAA GAG GAG ACA GTT AAT AAA GAT GTT AGA ATA TTT CTG TAA CTA AGT TTT CTC CTC TGT CAA TTA TTT CTA CAA S Y K D I D S K E E T V N K D V>

B-31 OSP / B-31 P-93

	1160				1	170			118	30		1:	1200			
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											GAC					
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	N	L	Q	К	T	K	P	Q	V	ĸ	D	Q	v	T	S	L>
			121	0		12	220		1	1230			124	10		
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٠.	T44	GAA	GAT	TTG	ACT	ACT	ATG	TCT	ATA	GAT	ŤCC	AGT	AGT	CCT	CTA	TTT
	TTA	CTT	CTA	AAC	TGA	TGA	TAC	AGA	TAT	CTA	AGG	TCA	TCA	GGA	CAT	212
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		1350 13														
	:	1350			13	60		1	370			1380			139	90
	:	1350		•	13	60	•	1	370			1380		•	139	90
	• CAG	CGG	TAT		TTA	TAT		CGT	• GAA	AAA	• GAT	TTG			ATT	AAA
	• CAG	CGG	TAT		TTA	TAT		CGT	• GAA	AAA	•	TTG			ATT	AAA
	• CAG GTC	CGG GCC	TAT ATA	CCT	TTA AAT	TAT ATA	CTT	CGT GCA	GAA CTT	AAA	• GAT	TTG AAC	CAA	CAA	ATT AAT	AAA TTT
	• CAG GTC	CGG GCC R	TAT ATA	CCT	ATT TAA I	TAT ATA	CTT E	CGT GCA R	GAA CTT E	AAA TTT K	GAT CTA D	TTG AAC L	CAA V	CAA V	ATT AAT I	AAA TTT K>
	• CAG GTC	CGG GCC R	TAT ATA Y	CCT	ATT TAA I	TAT ATA Y	CTT E	CGT GCA	GAA CTT E	AAA TTT K	GAT CTA D	TTG AAC	CAA V	CAA V	ATT AAT	AAA TTT K>
	CAG GTC Q	CGG GCC R 1	TAT ATA Y 400	CCT G	TTA AAT I	TAT ATA Y 1410	CTT E AAG	CGT GCA R	GAA CTT E 14 CAG	AAA TTT K 20	GAT CTA D	TTG AAC L 1	CAA V 430	CAA V	ATT TAA I	AAA TTT K> 1440
	CAG GTC Q	CGG GCC R 1 GAT CTA	TAT ATA Y 400 TCA	CCT G GGA CCT	ATT TAA I AAA	TAT ATA Y 1410 GCT CGA	E AAG	CGT GCA R CTT GAA	GAA CTT E 14 CAG	AAA TTT K 20 ATA TAT	GAT CTA D CTT GAA	TTG AAC L 1- GAT CTA	CAA V 430 AAA TTT	CAA V CTT GAA	ATT TAA I GAA CTT	AAA TTT K> 1440 AAT
	CAG GTC Q	CGG GCC R 1 GAT CTA	TAT ATA Y 400 TCA	CCT G GGA CCT	ATT TAA I AAA	TAT ATA Y 1410 GCT CGA	E AAG	CGT GCA R CTT GAA	GAA CTT E 14 CAG	AAA TTT K 20 ATA TAT	GAT CTA D	TTG AAC L 1- GAT CTA	CAA V 430 AAA TTT	CAA V CTT GAA	ATT TAA I GAA CTT	AAA TTT K> 1440 AAT
	CAG GTC Q ATG	CGG GCC R 1 GAT CTA	TAT ATA Y 400 TCA AGT S	CCT G GGA CCT	ATT TAA I AAA	TAT ATA Y 1410 GCT CGA A	E AAG	CGT GCA R CTT GAA L	GAA CTT E 14 CAG GTC	AAA TTT K 20 ATA TAT	GAT CTA D CTT GAA L	TTG AAC L 1- GAT CTA	CAA V 430 AAA TTT	CAA V CTT GAA L	ATT TAA I GAA CTT	AAA TTT K> 1440 AAT
	CAG GTC Q ATG TAC M	CGG GCC R 1 GAT CTA D	TAT ATA Y 400 TCA AGT S	GGA CCT G	ATT TAA I AAA TTT K	TAT ATA Y 1410 • GCT CGA A	AAG TTC K	CGT GCA R CTT GAA L	GAA CTT E 14 CAG GTC Q	AAA TTT K 20 ATA TAT I	GAT CTA D CTT GAA L	TTG AAC L 1. GAT CTA D	CAA V 430 AAA TTT K	CAA V CTT GAA L	ATT TAA I GAA CTT E	AAA TTT K> 144C AAT TTA N>
	CAG GTC Q ATG TAC M	CGG GCC R 1 GAT CTA D	TAT ATA Y 400 TCA AGT S	GGA CCT G GS0 GTA	ATT TAA I AAA TTT K	TAT ATA Y 1410 GCT CGA A 1	AAG TTC K	CGT GCA R CTT GAA L	GAA CTT E 14 CAG GTC Q	AAA TTT K 20 ATA TAT I 1470	GAT CTA D CTT GAA L	TTG AAC L 1- GAT CTA D	CAA V 430 AAA TTT K 14	CAA V CTT GAA L 80	ATT TAA I GAA CIT E	AAA TTT K> 1440 AAT TTA N>
	CAG GTC Q ATG TAC M	CGG GCC R 1 GAT CTA D	TAT ATA Y 400 TCA AGT S 14	GGA	ATT TAA I AAA TTT K TCA	TAT ATA Y 1410 GCT CGA A 1 GAG	AAG TTC K	CGT GCA R CTT GAA L	GAA CTT E 14 CAG GTC Q	AAA TTT K 20 ATA TAT I 1470	GAT CTA D CTT GAA L	TTG AAC L 1- GAT CTA D AAT TTA	CAA V 430 AAA TTT K 14 AAA TTT	CAA V CTT GAA L 80	ATT TAA I GAA CTT E TCA AGT	AAA TTT K> 1440 AAT TTA N>
	CAG GTC Q ATG TAC M	CGG GCC R 1 GAT CTA D	TAT ATA Y 400 TCA AGT S 14	GGA	ATT TAA I AAA TTT K TCA	TAT ATA Y 1410 GCT CGA A 1 GAG	AAG TTC K	CGT GCA R CTT GAA L	GAA CTT E 14 CAG GTC Q	AAA TTT K 20 ATA TAT I 1470	GAT CTA D CTT GAA L	TTG AAC L 1- GAT CTA D AAT TTA	CAA V 430 AAA TTT K 14 AAA TTT	CAA V CTT GAA L 80	ATT TAA I GAA CTT E TCA AGT	AAA TTT K> 1440 AAT TTA N>
1	CAG GTC Q ATG TAC M	CGG GCC R 1 GAT CTA D	TAT ATA Y 400 TCA AGT S 14	GGA	ATT TAA I AAA TTT K TCA AGT S	TAT ATA Y 1410 GCT CGA A 1 GAG	AAG TTC K 460 TCT AGA	CGT GCA R CTT GAA L	GAA CTT E 14 CAG GTC Q	AAA TTT K 20 ATA TAT I 1470 GAG	GAT CTA D CTT GAA L	TTG AAC L 1. GAT CTA D AAT TTA	CAA V 430 AAA TTT K 14 AAA TTT K	CAA V CTT GAA L 80	ATT TAA I GAA CTT E TCA AGT S	AAA TTT K> 1446 AAT TTA N>
1	CAG GTC Q ATG TAC M	CGG GCC R 1 GAT CTA D	TATA Y 400 TCA AGT S 14	GGA GGA CCT G SO CA1 V	ATT TAA I AAA TTT K TCA TCA	TAT ATA Y 1410 GCT CGA A 1 GAG CTC	AAG TTC K 460 TCT AGA S	CGT GCA R CTT GAA L AAT TTA N	GAA CTT E 14 CAG GTC Q	AAA TTT K 20 ATA TAT I 1470 GAG CTC	GAT CTA D CTT GAA L ATT TAA I	TTG AAC L 1. GAT CTA D AAT TTA	CAA V 430 AAA TIT K AAA TIT K	CAA V CTT GAA L 80 AAT TTA N	ATT TAA I GAA CTT E TCA AGT S	AAA TTT K> 1446 AAT TTA N>
1	CAG GTC Q ATG TAC M TTA AAT L	CGG GCC R 1 GAT CTA D	TATA Y 400 TCA AGT S 14 CGT CAC V	GGA GGA GGA GGA GGA CA1 V	ATT TAA I AAA TTT K TCA TCA	TAT ATA Y 1410 . GCT CGA A 1 . GAG	AAG TTC K 460 . TCT AGA S 15	CGT GCA R CTT GAA L AAT N	GAA CTT E 14 CAG GTC Q	AAA TTT K 20 ATA TAT I 1470 GAG CTC	GAT CTA D CTT GAA L ATT TAA I	TTG AAC L 1. GAT CTA D AAT TTA N	CAA V 430 AAA TIT K AAA TIT K	CAA V CTT GAA L 80 AAT TTA N	ATT TAA I GAA CTT E TCA AGT S	AAA TTT K> 144C AAT TTA N>

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71/133

B-31 OSP A/ B-31 P-93

1540 1550 1560 1570 AGT AGT AAT GAT TGG AGA TTG GCC AAA TTT TCT CCT AAA AAT TTA GAT TCA TCA TTA CTA ACC TCT AAC CGG TTT AAA AGA GGA TTT TTA AAT CTA S S N D W R L A K F S P K N L D> 1590 1600 1610 1620 GAG TTT ATT CTT TCA GAG AAT AAA ATT ATG CCT TTT ACT AGC TTT TCT CTC AAA TAA GAA AGT CTC TTA TTT TAA TAC GGA_AAA, TGA TCG AAA AGA EFILSENKIM PFTS FS 1650 1640 1660 1670 • GTG AGA AAA AAT TTT ATT TAT TTG CAA GAT GAG TTT AAA AGT CTA GTT CAC TCT TTT TTA AAA TAA ATA AAC GTT CTA CTC AAA TTT TCA GAT CAA -V R K N F I Y L Q D E F K S L V> 1690 1700 1710 ATT TTA GAT GTA AAT ACT TTA AAA AAA GTT AAG GGT CAC C TAA AAT CTA CAT TTA TGA AAT TTT TTT CAA TTC CCA GTG G I L D V N T L K K V K G H X>

#### B-31 OSP B/ B-31 P41 (122-234)

OSPB/Fla122-234

Sequence Range: 1 to 1180

20 GCA CAA AAA GGT GCT GAG TCA ATT GGT TCT CAA AAA GAA AAT GAT CTA CGT GTT TTT CCA CGA CTC AGT TAA CCA AGA GTT TTT CTT TTA CTA GAT AQKGAESIGSQKENDL> _ 80 ____. 60 70 AAC CTT GAA GAC TCT AGT AAA AAA TCA CAT CAA AAC GCT AAA CAA GAC TTG GAA CTT CTG AGA TCA TTT TTT AGT GTA GTT TTG CGA TTT GTT CTG N L E D S S K K S H Q N A K Q D> 110 120 130 CTT CCT GCG GTG ACA GAA GAC TCA GTG TCT TTG TTT AAT GGT AAT AAA GAA GGA CGC CAC TGT CTT CTG AGT CAC AGA AAC AAA TTA CCA TTA TTT L P A V T E D S V S L F N G N K> 160 170 180 ATT TTT GTA AGC AAA GAA AAA AAT AGC TCC GGC AAA TAT GAT TTA AGA TAA AAA CAT TCG TTT CTT TTT TTA TCG AGG CCG TTT ATA CTA AAT TCT I F V S K E K N S S G K Y D. L R> 200 210 220 230 GCA ACA ATT GAT CAG GTT GAA CTT AAA GGA ACT TCC GAT AAA AAC AAT CGT TGT TAA CTA GTC CAA CTT GAA TTT CCT TGA AGG CTA TTT TTG TTA A T I D Q V E L K G T S D K N N> 260 270 250 GGT TCT GGA ACC CTT GAA GGT TCA AAG CCT GAC AAG AGT AAA GTA AAA CCA AGA CCT TGG GAA CTT CCA AGT TTC GGA CTG TTC TCA TTT CAT TTT G S G T L E G S K P D K S K V K> 300 310 ·320 330 TTA ACA GTT TCT GCT GAT TTA AAC ACA GTA ACC TTA GAA GCA TTT GAT AAT TGT CAA AGA CGA CTA AAT TTG TGT CAT TGG AAT CTT CGT AAA CTA L T V S A D L N T V T L E A . F D> 340 350 360 370 380 GCC AGC AAC CAA AAA ATT TCA AGT AAA GTT ACT AAA AAA CAG GGG TCA CGG TCG TTG GTT TTT TAA AGT TCA TTT CAA TGA TTT TTT GTC CCC AGT A S N Q K I S S K V T K K Q G S>

### B-31 OSP B/ B-31 P41 (122-234)

		400				410				420		430						
	•	•		•		•	•		•		•	•		•		•		
	ATA	ACA	GAG	GAA	ACT	CTC	AAA	GCT	TAA	AAA	TTA	GAC	TCA	AAG	AAA	TTA		
	TAT	TGT	CTC	CTT	TGA	GAG	TTT	CGA	TTA	TTT	TAA	CTG	AGT	TTC	TTT	TAA		
	I	T	•E .	E	T	L	K	A	N	K	L	D	S	K.	K	L>		
		4	140			450			40	<b>50</b>		4	170			480		
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ŗ.	ACA	AGA	TCA	AAC	GGA	ACT	ACA	CTT	GAA	TAC	TCA	CAA	ATA	ACA	GAT	شک		
	TGT	TCT	AGT	TTG	CCT	TGA	TGT	GAA	CTT	ATG	AGT	GTT	TAT	TGT	CTA	CGA		
	T	R	S	N	G	T	T	L	E	Υ	`s¯		I			۸>		
			49	90		500				510			c.	20				
		•	٠.	•	•	•	•		•	210	-	•	34	20		٠.		
	GAC	AAT	GCT	ACA	AAA	GCA	GTA	GAA	ACT	СТА	444	таа	ACC.	እ <b>ጥ</b> ጥ		~~~		
	CTG	TTA	CGA	TGT	TTT	CGT	CAT	CTT	TGA	GAT	TTT	TTA	TCG	TAA	TTC	CIA		
	D	N	A	T	K	A	V	E	T	L	к	N	-5	Ť	TIC.	٠.٠.		
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	530			540			55	0		9	60		• ·	570				
	•		•	•		•		•	•		•		•	•		•		
	GAA	GGA	AGT	CTT	GTA	GTC	GGA	AAA	ÄCA	ACA	GTG	GAA	TTA	AAA	GAA	GGT		
				GAA	CAT	CAG	CCT	TTT	TGT	TGT	CAC		TAA	TTT	CTT	CCA		
	£	G	S	L	V	V	G	K	T	T	V	Ε	I	K	E	G>		
	580				590				600 6				510			620		
	58	30		9	90			600			61	.0		6	20			
	_	*	•		•			•		•		•	•		•			
	ACT	• GTT	ACT	CTA	• AAA	AGA	• GAA	TTA	GAA	• AAA	GAT	• GGA	AAA	GTA	AAA	GTC		
	ACT	• GTT	ACT TGA		AAA TTT	TCT	CTT	TTA	CTT	AAA TTT	GAT	• GGA	AAA TTT	GTA	AAA	GTC CAG		
	ACT TGA	• GTT	ACT TGA T	CTA	AAA TTT	AGA TCT R	CTT	TTA	CTT	TTT	GAT	• GGA CCT	AAA TTT K	GTA	AAA TTT	GTC CAG V>		
	ACT TGA	GTT CAA V	TGA	CTA GAT	AAA TTT K	TCT R	CTT	ATT TAA I	CTT E	TTT	GAT CTA	• GGA CCT G	TTT	GTA CAT	AAA TTT K	CAG V>		
	ACT TGA	GTT CAA	TGA	CTA GAT	AAA TTT	TCT R	CTT	ATT TAA I	CTT	TTT	GAT CTA	• GGA CCT	TTT	GTA CAT	AAA TTT	CAG V>		
	ACT TGA T	GTT CAA V	TGA T	CTA GAT L	AAA TTT K	TCT R	CTT E	ATT TAA I	E E 550	TTT K	GAT CTA D	GGA CCT G	TTT K	GTA CAT V	AAA TTT K	CAG V>		
	ACT TGA T	GTT CAA V 630	TGA T AAT	CTA GAT L GAC CTG	AAA TTT K 64 ACT TGA	TCT R 10 • GCA CGT	E GGT CCA	ATT TAA I TCT AGA	E 550 AAC TTG	TTT K AAA TTT	GAT CTA D	GGA CCT G 660	TTT K GGT	GTA CAT V	AAA TTT K 67	CAG V> 0		
	ACT TGA T	GTT CAA V 630	TGA T AAT TTA	CTA GAT L GAC CTG	AAA TTT K 64 ACT TGA	TCT R 10 • GCA	E GGT CCA	ATT TAA I TCT AGA	E 550 AAC TTG	TTT K AAA TTT	GAT CTA D	GGA CCT G 660 ACA TGT	TTT K GGT	GTA CAT V • AAA TTT	AAA TTT K 67	CAG V> 0 GAA CTT		
	ACT TGA T	GTT CAA V 630 TTG AAC L	TGA T AAT TTA	CTA GAT L GAC CTG	AAA TTT K 64 ACT TGA	TCT R 10 • GCA CGT	E GGT CCA	ATT TAA I TCT AGA	E 550 AAC TTG N	TTT K AAA TTT K	GAT CTA D	GGA CCT G 660 ACA TGT T	TTT K GGT CCA G	GTA CAT V • AAA TTT	AAA TTT K 67 TGG ACC	CAG V> 0 GAA CTT E>		
	ACT TGA T	GTT CAA V 630 TTG AAC L	TGA T AAT TTA N	CTA GAT L GAC CTG	AAA TTT K 64 ACT TGA	TCT R 10 GCA CGT A	E GGT CCA	ATT TAA I TCT AGA	E 550 AAC TTG	TTT K AAA TTT K	GAT CTA D	GGA CCT G 660 ACA TGT T	TTT K GGT CCA	GTA CAT V • AAA TTT	AAA TTT K 67 TGG ACC	CAG V> 0 GAA CTT		
	ACT TGA T TTT AAA F	GTT CAA V 630 TTG AAC L	TGA T AAT TTA N	CTA GAT L GAC CTG D	AAA TTT K 64 ACT TGA T	TCT R 10 GCA CGT A 690	CTT E GGT CCA G	ATT TAA I TCT AGA S	E 550 AAC TTG N	AAA TTT K	GAT CTA D • AAA TTT K	GGA CCT G 660 ACA TGT T	GGT CCA G	GTA CAT V • AAA TTT K	AAA TTT K 67 TGG ACC W	CAG V>  0 GAA CTT E> 720		
	ACT TGA T TTT AAA F	GTT CAA V 630 AAC L AGT TCA	TGA T  AAT TTA N  80 ACT TGA	CTA GAT L GAC CTG D	AAA TTT K 64 ACT TGA T ACT TGA	TCT R 10 GCA CGT A 690 TTA AAT	CTT E GGT CCA G	ATT TAA I TCT AGA S ATT TAA	ETT E 550 AAC TTG N 70 AGT TCA	AAA TTT K	GAT CTA D AAA TTT K	GGA CCT G 660 ACA TGT T	GGT CCA G	GTA CAT V  AAA TTT K	AAA TTT K 67 TGG ACC W ACT TGA	CAG V>  0 * GAA CTT E> 720 AAA		
	ACT TGA T TTT AAA F	GTT CAA V 630 AAC L AGT TCA	TGA T  AAT TTA N  80 ACT TGA	CTA GAT L GAC CTG D	AAA TTT K 64 ACT TGA T ACT TGA	TCT R 10 GCA CGT A 690 TTA AAT	CTT E GGT CCA G	ATT TAA I TCT AGA S ATT TAA	ETT E 550 AAC TTG N 70 AGT TCA	AAA TTT K	GAT CTA D AAA TTT K	GGA CCT G 660 ACA TGT T	GGT CCA G	GTA CAT V  AAA TTT K	AAA TTT K 67 TGG ACC W ACT TGA	CAG V>  0 * GAA CTT E> 720 AAA		
	ACT TGA T TTT AAA F	GTT CAA V 630 AAC L AGT TCA	TGA T AAT TTA N 580 ACT TGA T	CTA GAT L GAC CTG D AGC TCG S	AAA TTT K 64 ACT TGA T ACT TGA	TCT R IO GCA CGT A 690 TTA AAT L	GGT CCA G ACA TGT	ATT TAA I TCT AGA S ATT TAA	ETT E 550 AAC TTG N 70 AGT TCA	AAA TTT K O GCT CGA A	GAT CTA D AAA TTT K	GGA CCT G 660 ACA TGT T	GGT CCA G 10 . AAA TTT K	GTA CAT V AAA TTT K AAA TTT K	AAA TTT K 67 TGG ACC W ACT TGA	CAG V> 0 *GAA CTT E> 720 AAA TTT		
	ACT TGA T TTT AAA F	GTT CAA V 630 AAC L AGT TCA	TGA T AAT TTA N 580 ACT TGA T	CTA GAT L GAC CTG D	AAA TTT K 64 ACT TGA T ACT TGA	TCT R IO GCA CGT A 690 TTA AAT L	CTT E GGT CCA G	ATT TAA I TCT AGA S ATT TAA	ETT E 550 AAC TTG N 70 AGT TCA	AAA TTT K	GAT CTA D AAA TTT K	GGA CCT G 660 ACA TGT T	GGT CCA G 10 . AAA TTT K	GTA CAT V  AAA TTT K	AAA TTT K 67 TGG ACC W ACT TGA	CAG V> 0 *GAA CTT E> 720 AAA TTT		
	ACT TGA T TTT AAA F GAC CTG	GTT CAA V 630 TTG AAC L AGT TCA S	TGA T AAT TTA N 580 ACT TGA T	CTA GAT L GAC CTG D AGC TCG S	AAA TTT K 64 ACT TGA T ACT TGA T	TCT R GCA CGT A 690 TTA AAT L	GGT CCA G ACA TGT T	ATT TAA I TCT AGA S ATT TAA I	ETT E 550 AAC TTG N 70 AGT TCA S	AAA TTT K O GCT CGA A	GAT CTA D AAA TTT K GAC CTG D	GGA CCT G 660 ACA TGT T	GGT CCA G 10 AAA TTT K	GTA CAT V  AAA TTT K  AAA TTT K	AAA TTT K 67 TGG ACC W ACT TGA T	CAG V> 0 * GAA CTT E> 720 * AAA TTT K>		
	ACT TGA T TTT AAA F GAC CTG D	GTT CAA V 630 TTG AAC L AGT TCA S	TGA T AAT TTA N 580 ACT TGA T	CTA GAT L GAC CTG D AGC TCG S	AAA TTT K 64 ACT TGA T ACT TGA T	TCT R IO GCA CGT A 690 TTA AAT L	GGT CCA G ACA TGT T	ATT TAA I TCT AGA S ATT TAA I	ETT E S50 AAC TTG N 70 AGT TCA S	AAA TTT K O GCT CGA A 750	GAT CTA D AAA TTT K GAC CTG D	GGA CCT G 660 ACA TGT T AGC TCG S	GGT CCA G 10 AAA TTT K 76 CAA	GTA CAT V AAA TTT K AAA TTT K CAA	AAA TTT K 67 TGG ACC W ACT TGA T TAC	CAG V> 0 * GAA CTT E> 720 * AAA TTT K>		

FIGURE 33 (2 of 4)

B-31 OSP B/ B-31 P41 (122-234)

810 770 780 790 • ACA GCT GGA ACC AGC CTA GAA GGA TCA GCA AGT GAA ATT AAA AAT CTT TGT CGA CCT TGG TCG GAT CTT CCT AGT CGT TCA CTT TAA TTT TTA GAA TAGTSLEGSAS.EIKN L 830 840 850 TCA GAG CTT AAA AAC GCT TTA AAA GGT CAC CCC ATG GCT CAA TAT AAC AGT CTC GAA TTT TTG CGA AAT TTT CCA GTG GGG TAC CGA GTT ATA TTG S E L K N A L K G H: P==M: A Q Y N> 870 880 890 890 900 CAA ATG CAC ATG TTA TCA AAC AAA TCT GCT TCT CAA AAT GTA AGA ACA GTT TAC GTG TAC AAT AGT TTG TTT AGA CGA AGA GTT TTA CAT TCT TGT Q M H M L S N K S A S Q N V R T> 920 930 940 950 , 960 GCT GAA GAG CTT GGA ATG CAG CCT GCA AAA ATT AAC ACA CCA GCA TCA CGA CTT CTC GAA CCT TAC GTC GGA CGT TTT TAA TTG TGT GGT CGT AGT A E E L G M Q P A K I N T P A S> 980 990 CTT TCA GGG CTT CAA GCG TCT TGG ACT TTA AGA GTT CAT GTT GGA GCA GAA AGT CCC GAA GTT CGC AGA ACC TGA AAT TCT CAA GTA CAA CCT CGT L S G L Q A S W T L R V H V G A> 1030 1040 1020 ACC CAA GAT GAA GCT ATT GCT GTA AAT ATT TAT GCA GCT AAT GTT GCA TGG GTT CTA CTT CGA TAA CGA CAT TTA TAA ATA CGT CGA TTA CAA CGT T Q D E A I A V N I Y A A N V A> 1070 1080 1090 1100 AAT CTT TTC TCT GGT GAG GGA GCT CAA ACT GCT CAG GCT GCA CCG GTT TTA GAA AAG AGA CCA CTC CCT CGA GTT TGA CGA GTC CGA CGT GGC CAA N L F S G E G A Q T A Q A A P V>1120 1130 1140 1110 CAA GAG GGT GTT CAA CAG GAA GGA GCT CAA CAG CCA GCA CCT GCT ACA GTT CTC CCA CAA GTT GTC CTT CCT CGA GTT GTC GGT CGT GGA CGA TGT Q E G V Q Q E G A Q Q P A P A T>

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75/133

B-31 OSP B/ B-31 P41 (122-234)

GCA CCT TCT CAA GGC GGA GTT GGT CAC C
CGT GGA AGA GTT CCG CCT CAA CCA GTG G
A P S Q G G V G H X>

FIGURE 33 (4 of 4)

B-31 OSP B / B-31 P41 (122-295)

Sequence Range: 1 to 1363

20 30 GCA CAA AAA GGT GCT GAG TCA ATT GGT TCT CAA AAA GAA AAT GAT CTA CGT GTT TTT CCA CGA CTC AGT TAA CCA AGA GTT TTT CTT TTA CTA GAT A Q K G A E S I G S Q K E N D L> 70 60 50 80 AAC CTT GAA GAC TCT AGT AAA AAA TCA CAT CAA-AAC GCT ALA CAA GAC TTG GAA CTT CTG AGA TCA TTT TTT AGT GTA GTT TTG CGA TTT GTT CTG N L E D S S K K S H Q N A K Q D> 100 110 120 130 140 CTT CCT GCG GTG ACA GAA GAC TCA GTG TCT TTG TTT AAT GGT AAT ALA GAA GGA CGC CAC TGT CTT CTG AGT CAC AGA AAC AAA TTA CCA TTA TTT L P A V T E D S V S L F N G N K> 150 160 170 180 ATT TTT GTA AGC AAA GAA AAA AAT AGC TCC GGC AAA TAT GAT TTA AGA TAA AAA CAT TCG TTT CTT TTT TTA TCG AGG CCG TTT ATA CTA AAT TCT I F V S K E K N S S G K Y D L R> 200 210 220 GCA ACA ATT GAT CAG GTT GAA CTT AAA GGA ACT TCC GAT AAA AAC AAT CGT TGT TAA CTA GTC CAA CTT GAA TTT CCT TGA AGG CTA TTT TTG TTA A T I D Q V E L K G T S D K N N> 250 260 270 GGT TCT GGA ACC CTT GAA GGT TCA AAG CCT GAC AAG AGT AAA GTA AAA CCA AGA CCT TGG GAA CTT CCA AGT TTC GGA CTG TTC TCA TTT CAT TTT G S G T L E G S K P D K S K V K> 300 310 320 . • TTA ACA GTT TCT GCT GAT TTA AAC ACA GTA ACC TTA GAA GCA TTT GAT AAT TGT CAA AGA CGA CTA AAT TTG TGT CAT TGG AAT CTT CGT AAA CTA L T V S A D L N T V T L E A F D> 350 360 370 340 GCC AGC AAC CAA AAA ATT TCA AGT AAA GTT ACT AAA AAA CAG GGG TCA CGG TCG TTG GTT TTT TAA AGT TCA TTT CAA TGA TTT TTT GTC CCC AGT A S N Q K I S S K V T K K Q G S>

FIGURE 34 (1 of 4)

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77/133

#### B-31 OSP B / B-31 P41 (122-295)

		390			4	00			410			420			4	30
	•	•		•		•	•		•		•	•		•		•
	ATA	ACA	GAG	GAA	ACT	CTC	AAA	GCT	AAT	AAA	TTA	GAC	TCA	AAG	AAA	TTA
	TAT	TGT	CTC	CTT	TGA	GAG	TTT	CGA	TTA	TTT	AAT	CTG	AGT	TTC	TTT	TAS
	I	T	E	E	T	L	ĸ	λ			L			ĸ		L>
															••	
		4	140			450			46	50		4	170			480
٠,	•		•		•	•		•		•	•		•		•	•
	ACA	AGA	TCA	AAC	GGA	ACT	ACA	CTT	Gλλ	TAC	TCA	CAA	ATA	ACA	GAT	GCT
	TGT	TCT	AGT	TTG	CCT	TGA	TGT	GAA	CTT	ATG-	AGT	GTT	TAT	TGT	CTA	CGA
	T	R		N		· T	T	L	E	Y	S	Q	I	T	D	λ>
			49	90		5	500			510			52	20	•	٠.
		*		•	•		•		•	*		•		•	•	
	GAC	AAT	GCT	ACA	AAA	GCA	GTA	GAA	ACT	CTA	AAA	AAT	AGC	TTA	ÄAG	CTT
	CTG	TTA	CGA	TGT	TTT	CGT	CAT	CTT	TGA	GAT	TTT	TTA	TCG	TAA	TTC	GAA
	Ð	N	Α	T	K	A	V	E	T	L	K	N	S	I	K	L>
	530			C 4 0			55	. ^							•	
	220		•	540			٥.	•		-	60			570		
	G) k	CCA	ACT	СТТ	GTA	CTC	CCA	**	NCX.	NCN.	CTC	C	****	•		•
	CTT	CCT	TCA	CII	CAT	CYC	CCT	TTT	TOT	TCT	CVC	CTT	WII	AAA	GAA	GGT
	E	G	S	L	V	V	G	K	T	T	V	E				
	-	J			•	•	•	• • • • • • • • • • • • • • • • • • • •	4	•	•	L	I	K	£	G>
	580			9	590			600			61	0		620		
		•	•		•		•	•		•		•	•		•	
	ACT	GTT	ACT	CTA	AAA	AGA	GAA	TTA	GAA	AAA	GAT	GGA	AAA	GTA	AAA	GTC
	TGA	CAA	TGA	GAT	TTT	TCT	CTT	AAT	CTT	TTT	CTA	CCT	TTT	CAT	TTT	CAG
	T	V	T	. L	K	R	E	I	Ε	ĸ	D	G	K	v	K	V>
		630			64	0		•	550			660			67	0
	•	•		•		•	•		•		•	•		•		•
					ACT											
					TGA											
	F	L	И	D	T	A	G	S	N	K	K	T	G	K	W	E>
		690			700			١0		+	10	720				
	•	•	580 •		•	•		•	•	•	*	•	10		•	120
	GAC	AGT	ACT	AGC	ACT	TTA	ACA	ATT	AGT	GCT	GAC	AGC	444	AAA	<b>∆</b> CT	222
					TGA											
	D	s	T	s	T	L	T	I	S	A	D	S	к	ĸ	T	K>
											i				_	•
			7	30		-	740 75			750			76	50		
		•		•	•		•		•	•		•		• .	•	
					TTA											
	CTA	220	CAC	AAC	TAA	TCT	AT)	CCA	TCT	TAA	TGT	CAT	CTT	$\sim$	8 m~	<b>~~~</b>
					7 U. 1										MIG	110

FIGURE 34 (2 of 4)

. . . .

B-31 OSP B / B-31 P41 (122-295)

770 780 790 800 810 ACA GCT GGA ACC AGC CTA GAA GGA TCA GCA AGT GAA ATT AAA AAT CTT TGT CGA CCT TGG TCG GAT CTT CCT AGT CGT TCA CTT TAA TIT TTA GAA TAGTSLEGSASEIKN L> 850 820 830 840 860 • TCA GAG CTT AAA AAC GCT TTA AAA GGT CAC CCC ATG GCT CAA TAT AAC AGT CTC GAA TTT TTG CGA AAT TTT CCA GTG GGG TAC CGA GTT ATA TTG S E L K N A L K G H P M A Q Y N> 880 890 900 870 CAA ATG CAC ATG TTA TCA AAC AAA TCT GCT TCT CAA AAT GTA AGA ACA GTT TAC GTG TAC AAT AGT TTG TTT AGA CGA AGA GTT TTA CAT TCT TGT Q M H M L S N K S A S Q N V R T> 920 930 940 950 GCT GAA GAG CTT GGA ATG CAG CCT GCA AAA ATT AAC ACA CCA GCA TCA CGA CTT CTC GAA CCT TAC GTC GGA CGT TTT TAA TTG TGT GGT CGT AGT A E E L G M Q P A K I N T P A S> 970 980 990 1000 CTT TCA GGG CTT CAA GCG TCT TGG ACT TTA AGA GTT CAT GTT GGA GCA GAA AGT CCC GAA GTT CGC AGA ACC TGA AAT TCT CAA GTA CAA CCT CGT L S G L Q A S W T L R V H V G &> 1020 1030 1040 1010 1050 ACC CAA GAT GAA GCT ATT GCT GTA AAT ATT TAT GCA GCT AAT GTT GCA TGG GTT CTA CTT CGA TAA CGA CAT TTA TAA ATA CGT CGA TTA CAA CGT T Q D E A I A V N I Y A A N V A> 1060 1070 1080 1090 1100 AAT CTT TTC TCT GGT GAG GGA GCT CAA ACT GCT CAG GCT GCA CCG GTT TTA GAA AAG AGA CCA CTC CCT CGA GTT TGA CGA GTC CGA CGT GGC CAA N L F S G E G A Q T A Q A A P V> 1120 1130 1140 CAA GAG GGT GTT CAA CAG GAA GGA GCT CAA CAG CCA GCA CCT GCT ACA GTT CTC CCA CAA GTT GTC CTT CCT CGA GTT GTC GGT CGT GGA CGA TGT

FIGURE: 34 (3 of 4)

Q E G V Q Q E G A Q Q P A P A T>

#### B-31 OSP B / B-31 P41 (122-295)

1180 1190 1160 1170 GCA CCT TCT CAA GGC GGA GTT AAT TCT CCT GTT AAT GTT ACA ACT ACA CGT GGA AGA GTT CCG CCT CAA TTA AGA GGA CAA TTA CAA TGT TGA TGT A P S Q G G V N S P V N V T T T> 1230 ` 1220 1210 GTT GAT GCT AAT ACA TCA CTT GCT AAA ATT GAA AAT GCT ATT AGA ATG CAA CTA CGA TŢA TGT AGT GAA CGA TTT TAA CTT TTA CGA TAA TCT TAC V D A N T S L A K I E N A I R M> 1250 1260 1270 1280 . 1295 • ATA AGT GAT CAA AGG GCA AAT TTA GGT GCT TTC CAA AAT AGA CTT GAA TAT TCA CTA GTT TCC CGT TTA AAT CCA CGA AAG GTT TTA TCT GAA CTT I S D Q R A N L G A F Q N R L E> 1310 1320 1330 TCT ATA AAG AAT AGT ACT GAG TAT GCA ATT GAA AAT CTA AAA GCA TCT AGA TAT TTC TTA TCA TGA CTC ATA CGT TAA CTT TTA GAT TTT CGT AGA S I K N S T E Y A I E N L K A S> 1360 1350

TAT GCT CAA ATA GGT CAC C
ATA CGA GTT TAT CCA GTG G
Y A Q I G H X>

FIGURE 34 (4 of 4)

B-31 OSP B/ B-31 P41 (140-234)

Sequence Range: 1 to 1141

20 10 30 GCA CAA AAA GGT GCT GAG TCA ATT GGT TCT CAA AAA GAA AAT GAT CTA CGT GTT TTT CCA CGA CTC AGT TAA CCA AGA GTT TTT CTT TTA CTA GAT A Q K G A E S I G S Q K E N D L> 70 60 • AAC CTT GAA GAC TCT AGT AAA AAA TCA CAT CAA AAC GCT AAA CAA GAC TTG GAA CTT CTG AGA TCA TTT TTT AGT GTA GTT TTG CGA TTT GTT CTG N L E D S S K K S H Q N A K Q D> 100 110 120 130 • CTT CCT GCG GTG ACA GAA GAC TCA GTG TCT TTG TTT AAT GGT AAT AAA GAA GGA CGC CAC TGT CTT CTG AGT CAC AGA AAC AAA TTA CCA TTA TTT L P A V T E D S V S L F N G N K> 170 150 160 180 190 ATT TTT GTA AGC AAA GAA AAA AAT AGC TCC GGC AAA TAT GAT TTA AGA TAA AAA CAT TCG TTT CTT TTT TTA TCG AGG CCG TTT ATA CTA AAT TCT I F V S K E K N S S G K Y D L R> 230 210 220 200 GCA ACA ATT GAT CAG GTT GAA CTT AAA GGA ACT TCC GAT AAA AAC AAT CGT TGT TAA CTA GTC CAA CTT GAA TTT CCT TGA AGG CTA TTT TTG TTA A T I D Q V E L K G T S D K N N> 250 260 270 280 GGT TCT GGA ACC CTT GAA GGT TCA AAG CCT GAC AAG AGT AAA GTA AAA CCA AGA CCT TGG GAA CTT CCA AGT TTC GGA CTG TTC TCA TTT CAT TTT G S G T L E G S K P D K S K V K> 300 310 320 330 • TTA ACA GTT TCT GCT GAT TTA AAC ACA-GTA ACC TTA GAA GCA TTT GAT AAT TGT CAA AGA CGA CTA AAT TTG TGT CAT TGG AAT CTT CGT AAA CTA L T V S A D L N T V T L E A F DS 350 360 370 • GCC AGC AAC CAA AAA ATT TCA AGT AAA GTT ACT AAA AAA CAG GGG TCA CGG TCG TTG GTT TTT TAA AGT TCA TTT CAA TGA TTT TTT GTC CCC AGT A S N Q K I S S K V T K K Q G S>

### B-31 OSP B/ B-31 P41 (140-234)

				4.0	. ^			10			420			43	20
	390		•	40	•	•	•	*		•	420		•	٧.	•
ATA	ACA	GAG	GAA	ACT	СТС	AAA	GCT	TAA	AAA	ATT	GAC	TCA	AAG	AAA	TTA.
					GAG										
					L										
	4	40			450		_	46	50	_	4	170		_	480
. •		-		•	ACT		<u>-</u>	CNN	<b>ፕ</b> እር	TCN	CAA	እጥአ	A.C.N	Cam	-
					TGA										
					T										
•	1		.,	٥.	•	•	_	-		,	~~~ «	-	•	_	***
		49	90		5	00			510			52	20		
	•		•	•		•		•	•		•		•	. •.	
					GCA										
CTG					CGT										
D	N	A	T	ĸ	Α	V	E	T	L	K	N	. S	1	K	L>
530			540			55	50		•	560		•	570		
•		•	•		•	-	•	•	-	•		•	•		•
GAA	GGA	AGT	CTT	GTA	GTC	GGA	AAA	ACA	ACA	GTG	GAA	ATT	AAA	GAA	GGT
					CAG										
E	G	S	L	v	V	G	K	T	T	V	E	I	K	E	G>
<b>C</b> (	0.0			- 0 0			600			6	10			เวก	
58	B0 •	•		590		•	600		•	6:	10	•	•	520	
	•	• ACT		•	AGA	•	•	GAA	• AAA		•			•	GTC
ACT	• GŢT		CTA	• AAA	AGA TCT	• GAA	ATT			GAT	• GGA		GTA	• AAA	
ACT	• GŢT CAA		CTA GAT	AAA TTT		GAA CTT	ATT TAA	CTT	TTT	GAT CTA	• GGA CCT	TTT	GTA CAT	• AAA	CAG
ACT TGA	GȚT CAA V	TGA T	CTA GAT	AAA TTT K	TCT R	GAA CTT	ATT TAA I	CTT E.	TTT	GAT CTA D	GGA CCT G	TTT	GTA CAT	AAA TTT K	CAG V>
ACT TGA	• GŢT CAA	TGA T	CTA GAT	AAA TTT	TCT R	GAA CTT	ATT TAA I	CTT	TTT	GAT CTA D	• GGA CCT	TTT	GTA CAT	AAA TTT	CAG V>
ACT TGA T	GTT CAA V	TGA T	CTA GAT L	AAA TTT K	TCT R 40	GAA CTT E	ATT TAA I	CTT E.	TTT	GAT CTA D	GGA CCT G	TTT	GTA CAT V	AAA TTT K	CAG V>
ACT TGA T	GTT CAA V	TGA T	CTA GAT L	AAA TTT K 64	TCT R 40 • GCA	GAA CTT E	ATT TAA I TCT	CTT E. 650 AAC	TTT K	GAT CTA D	GGA CCT G 660 ACA	TTT K GGT	GTA CAT V	AAA TTT K 67	CAG V> 70 GAA
ACT TGA T TTT	GTT CAA V 630 TTG AAC	TGA T AAT TTA	CTA GAT L GAC CTG	AAA TTT K 64 ACT TGA	TCT R 40	GAA CTT E  GGT CCA	ATT TAA I TCT AGA	CTT E. 650 AAC TTG	TTT K AAA TTT	GAT CTA D	GGA CCT G 660 ACA TGT	TTT K GGT CCA	GTA CAT V AAA TTT	AAA TTT K 6' TGG ACC	CAG V> 70 GAA CTT
ACT TGA T TTT	GTT CAA V 630 TTG AAC	TGA T AAT TTA	CTA GAT L GAC CTG	AAA TTT K 64 ACT TGA	TCT R 40 • GCA CGT	GAA CTT E  GGT CCA	ATT TAA I TCT AGA	CTT E. 650 AAC TTG N	TTT K AAA TTT K	GAT CTA D	GGA CCT G 660 ACA TGT	TTT K GGT CCA G	GTA CAT V AAA TTT	AAA TTT K 6' TGG ACC	CAG V> 70 GAA CTT E>
ACT TGA T TTT	GTT CAA V 630 TTG AAC L	TGA T AAT TTA	CTA GAT L GAC CTG	AAA TTT K 64 ACT TGA	TCT R 40 • GCA CGT	GAA CTT E  GGT CCA	ATT TAA I TCT AGA	CTT E. 650 AAC TTG N	TTT K AAA TTT	GAT CTA D	GGA CCT G 660 ACA TGT	TTT K GGT CCA	GTA CAT V AAA TTT	AAA TTT K 6' TGG ACC	CAG V> 70 GAA CTT
ACT TGA T TTT AAA F	GTT CAA V 630 TTG AAC L	TGA T AAT TTA N 680	CTA GAT L GAC CTG	AAA TTT K 60 ACT TGA T	TCT R 40 GCA CGT A 690	GAA CTT E GGT CCA G	ATT TAA I TCT AGA S	CTT E. 650 AAC TTG N	TTT K AAA TTT K	GAT CTA D • AAA TTT K	GGA CCT G 660 ACA TGT T	GGT CCA G	GTA CAT V AAA TTT K	AAA TTT K 67 TGG ACC W	CAG V> 70 GAA CTT E> 720
ACT TGA T TTT AAA F	GTT CAA V 630 TTG AAC L	TGA T AAT TTA N 680 ACT	CTA GAT L GAC CTG D	AAA TTT K 6 ACT TGA T	TCT R 40 GCA CGT A 690	GAA CTT E GGT CCA G	ATT TAA I TCT AGA S	CTT E. 650 AAC TTG N 7	AAA TTT K  OO GCT	GAT CTA D AAA TTT K	GGA CCT G 660 ACA TGT T	GGT CCA G	GTA CAT V AAA TTT K	AAA TTT K 67 TGG ACC W	CAG V> 70 GAA CTT E> 720
ACT TGA T TTT AAA F GAC CTG	GTT CAA V 630 AAC L L AGT	TGA T AAT TTA N 680 ACT TGA	CTA GAT L GAC CTG D AGC	AAA TTT K 6 ACT TGA T ACT TGA	TCT R 40 GCA CGT A 690 TTA AAT	GAA CTT E GGT CCA G	ATT TAA I TCT AGA S ATT TAA	CTT E. 650 AAC TTG N 7 AGT	AAA TTT K  OO GCT	GAT CTA D AAA TTT K	GGA CCT G 660 ACA TGT T	GGT CCA G	GTA CAT V AAA TTT K	AAA TTT K 67 TGG ACC W ACT TGA	CAG V> 70 GAA CTT E> 720 AAA TTT
ACT TGA T TTT AAA F	GTT CAA V 630 AAC L L AGT	TGA T AAT TTA N 680 ACT TGA	CTA GAT L GAC CTG D AGC	AAA TTT K 6 ACT TGA T ACT TGA	TCT R 40 GCA CGT A 690	GAA CTT E GGT CCA G	ATT TAA I TCT AGA S ATT TAA	CTT E. 650 AAC TTG N 7 AGT	AAA TTT K  OO GCT	GAT CTA D AAA TTT K	GGA CCT G 660 ACA TGT T	GGT CCA G	GTA CAT V AAA TTT K	AAA TTT K 67 TGG ACC W ACT TGA	CAG V> 70 GAA CTT E> 720 AAA TTT
ACT TGA T TTT AAA F GAC CTG	GTT CAA V 630 AAC L L AGT	TGA T AAT TTA N 680 ACT TGA	CTA GAT L GAC CTG D AGC	AAA TTT K 6 ACT TGA T ACT TGA	TCT R 40 GCA CGT A 690 TTA AAT L	GAA CTT E GGT CCA G	ATT TAA I TCT AGA S ATT TAA	CTT E. 650 AAC TTG N 7 AGT	AAA TTT K  OO GCT	GAT CTA D AAA TTT K GAC CTG	GGA CCT G 660 ACA TGT T	GGT CCA G 710 AAA TTT K	GTA CAT V AAA TTT K	AAA TTT K 67 TGG ACC W ACT TGA	CAG V> 70 GAA CTT E> 720 AAA TTT
ACT TGA T TTT AAA F GAC CTG	GTT CAA V 630 TTG AAC L	TGA T AAT TTA N 680 TGA TGA	GAT L GAC CTG D AGC	AAA TTT K 6ACT TGA T ACT TGA T	TCT R 40 GCA CGT A 690 TTA AAT L	GAA CTT E GGT CCA G	ATT TAA I TCT AGA S ATT TAA I	CTT E 650 AAC TTG N 7 AGT TCA S	AAA TTT K  OO GCT CGA A 750	GAT CTA D AAA TTT K GAC CTG	GGA CCT G 660 ACA TGT T	GGT CCA G 710 . AAA TTT K	GTA CAT V  AAA TTT K  AAA TTT K	AAA TTT K 6° TGG ACC W ACT TGA	CAG V> 70 GAA CTT E> 720 AAA TTT K>
ACT TGA T TTT AAA F GAC CTG D	GTT CAA V 630 TTG AAC L	TGA T  AAT TTA N  680 TGA TGA TGA	GAT L GAC CTG D AGC	AAA TTT K 64 ACT TGA T ACT TGA T TTA	TCT R 40. GCA CGT A 690. TTA AAT L	GAA CTT E GGT CCA G ACA TGT T	ATT TAA I TCT AGA S ATT TAA I	CTT E. 650 AAC TTG N 7 AGT TCA S	AAA TTT K  00 GCT CGA 750	GAT CTA D AAA TTT K GAC CTG	GGA CCT G 660 ACA TGT T	GGT CCA G 710 . AAA TTT K 7	GTA CAT V  AAA TTT K  AAA TTT K	AAA TTT K 6T TGG ACC W ACT TGA T	CAG V> 70 GAA CTT E> 720 AAA TTT K>
ACT TGA T TTT AAA F GAC CTG D	GTT CAA V 630 AAC L AGT S	TGA T AAT TTA N 680 TGA TGA TGA TGGCCAC	GAC CTG D AGC S 30	AAA TTT K 64 ACT TGA T ACT TGA T TTA	TCT R 40. GCA CGT A 690. TTA AAT L	GAA CTT E GGT CCA G ACA TGT T	ATT TAA I TCT AGA S ATT TAA I	CTT E. 650 AAC TTG N 7 AGT TCA S	AAA TTT K  00 GCT CGA A 750	GAT CTA D AAA TTT K GAC CTG D	GGA CCT G 660 ACA TGT T AGC TCG S	GGT CCA G 710 . AAA TTT K 7	GTA CAT V  AAA TTT K  AAA TTT K	AAA TTT K 67 TGG ACC W ACT TGA T TAC	CAG V>  70 GAA CTT E>  720 AAA TTT K>  AAC TTG

FIGURE 35 (2 of 3)

#### B-31 OSP B/ B-31 P41 (140-234)

770 780 790 800 ACA GCT GGA ACC AGC CTA GAA GGA TCA GCA AGT GAA ATT AAA AAT CTT TGT CGA CCT TGG TCG GAT CTT CCT AGT CGT TCA CTT TAA TTT TTA GAA TAGTS LEGS ASEIKN L> 830 840 850 860 820 TCA GAG CTT AAA AAC GCT TTA AAA GGT CAC CCC ATG GCT TCT CAA AAT AGT CTC GAA TTT TTG CGA AAT TTT CCA GTG GGG TAC CGA AGA GTT TTA S E L K N A L K G H P M A S Q N> . 880 890 900 870 GTA AGA ACA GCT GAA GAG CTT GGA ATG CAG CCT GCA AAA ATT AAC ACA CAT TCT TGT CGA CTT CTC GAA CCT TAC GTC GGA CGT TTT TAA TTG TGT V R T A E E L G M Q P A K I N T> 920 930 940 950 CCA GCA TCA CTT TCA GGG CTT CAA GCG TCT TGG ACT TTA AGA GTT CAT GGT CGT AGT GAA AGT CCC GAA GTT CGC AGA ACC TGA AAT TCT CAA GTA PASLS GLQAS WTLRVH> 970 980 990 1000 • • GTT GGA GCA ACC CAA GAT GAA GCT ATT GCT GTA AAT ATT TAT GCA GCT CAA CCT CGT TGG GTT CTA CTT CGA TAA CGA CAT TTA TAA ATA CGT CGA V G A T O D E A I A V N I Y A A> 1020 1030 1040 1050 • AAT GTT GCA AAT CTT TTC TCT GGT GAG GGA GCT CAA ACT GCT CAG GCT TTA CAA CGT TTA GAA AAG AGA CCA CTC CCT CGA GTT TGA CGA GTC CGA NVANLFSGEGAQTAQA> 1070 1080 1090 GCA CCG GTT CAA GAG GGT GTT CAA CAG GAA GGA GCT CAA CAG CCA GCA CGT GGC CAA GTT CTC CCA CAA GTT GTC CTT CCT CGA GTT GTC GGT CGT A P V Q E G V Q Q E G A Q Q P A> 1120 1130 1140 CCT GCT ACA GCA CCT TCT CAA GGC GGA GTT GGT CAC C GGA CGA TGT CGT GGA AGA GTT CCG CCT CAA CCA GTG G PATAPS QGG VGH X>

FIGURE 35 (3 of 3)

#### B-31 OSP B/ B-31 P41 (140 -295)

Sequence Range: 1 to 1324

20 30 GCA CAA AAA GGT GCT GAG TCA ATT GGT TCT CAA AAA GAA AAT GAT CTA CGT GTT TTT CCA CGA CTC AGT TAA CCA AGA GTT TTT CTT TTA CTA GAT AQKGAESIGSQKEND L> 60 70 80 50 • · · · · · · AAC CTT GAA GAC TCT AGT AAA AAA TCA CAT CAA AAC GCT AAA CAA GAC TTG GAA CTT CTG AGA TCA TTT TTT AGT GTA GTT TTG CGA TTT GTT CTG N L E D S S K K S H Q N A K. Q. D> 120 130 100 110 CTT CCT GCG GTG ACA GAA GAC TCA GTG TCT TTG TTT - AAT GGT AAT AAA GAA GGA CGC CAC TGT CTT CTG AGT CAC AGA AAC AAA TTA CCA TTA TTT LPAVTEDSVSLFNGNK> 150 160 170 180 ATT TTT GTA AGC AAA GAA AAA AAT AGC TCC GGC AAA TAT GAT TTA AGA TAA AAA CAT TCG TTT CTT TTT TTA TCG AGG CCG TTT ATA CTA AAT TCT I F V S K E K N S S G K Y D L R> 200 210 220 230 GCA ACA ATT GAT CAG GTT GAA CTT AAA GGA ACT TCC GAT AAA AAC AAT CGT TGT TAA CTA GTC CAA CTT GAA TTT CCT TGA AGG CTA TTT TTG TTA ATIDQVELKGTSDKN N> 250 260 270 280 GGT TCT GGA ACC CTT GAA GGT TCA AAG CCT GAC AAG AGT AAA GTA AAA CCA AGA CCT TGG GAA CTT CCA AGT TTC GGA CTG TTC TCA TTT CAT TTT G S G T L E G S K P D K S K V K> 310 300 320 TTA ACA GTT TCT GCT GAT TTA AAC ACA GTA ACC TTA GAA GCA TTT GAT AAT TGT CAA AGA CGA CTA AAT TTG TGT CAT TGG AAT CTT CGT AAA CTA LTVSADLNTVTLEAFD> 370 380 350 360 GCC AGC AAC CAA AAA ATT TCA AGT AAA GTT ACT AAA AAA CAG GGG TCA CGG TCG TTG GTT TTT TAA AGT TCA TTT CAA TGA TTT TTT GTC CCC AGT ASNQKISSKVTKKQGS>

FIGURE 36 (1 of 4)

### B-31 OSP B/ B-31 P41 (140 -295)

		390			4	00			410			420	+		4	30
	•	•		•		•	•		•		•	•		•		•
	ATA	ACA	GAG	GAA	ACT	CTC	AAA	GCT	TAA	AAA	TTA	GAC	TCA	AAG	AAA	TTA
				CTT	TGA			CGA	TTA	TTT	AAT	CTG	AGT			
	Ι	T	E	E	T	L	K	A	N	K	L	D	S	K	K	L>
			440			450			4	60			470			480
•	. •		•		•	•		•		•	. •		•		•	•
	ACA	AGA	TCA	AAC	GGA	ACT	ACA	CTT	GAA	TAC	TCA	CAA	ATA	ACA	GAT	GCT
	T	R	AGT	TIG	CCT	TGA	TGT	GAA	CIT	ATG	AGT	GTT	TAT	TGT	CTA	CGA
	•	K	3	14	G	4	T	L	Ł	¥	- 5-	Q -	I	T	D	A>
			49	90		9	500			510			5	20		•
		•		•	•		•		•	•		•		•	• •	٠.
	GAC	AAT	GCT	ACA	AAA	GCA	GTA	GAA	ACT	CTA	AAA	AAT	AGC	ATT	AAG	CTT
	CTG	TTA	CGA	TGT	TTT	CGT	CAT	CII	TGA	GAT	TTT	TTA	TCG	TAA	TTC	GAA
	Ð	N	Α	T	K	A	V	E	T	L	K	N	Ş	I	K	L>
	530			540			55	50		9	560			570		
	•		•	•		•		•	•		•		•	•	,	•
	GAA	GGA	AGT	CTT	GTA	GTC	GGA	AAA	ACA	ACA	GTG	GAA	ATT	AAA	GAA	GGT
	CTT	CCT	TCA	GAA	CAT	CAG	CCT	TTT	TGT	TGT	CAC	CTT	TAA	TTT	CTT	CCA
	E	G	s	L	V	V	G	K	T	T	V	E	I	K	E	G>
	58	30		5	90			600			61	.0			520	
		•	•		•		•	•		•		•	•		•	
	ACT	GTT	ACT	CTA	AAA	AGA	GAA	TTA	GAA	AAA	GAT	GGλ	AAA	GTA	AAA	GTC
	T	V	1GY	GAT	TTT	TCT	CTT	TAA	CTT	TTT	CTA	CCI	TTT	CAT	TTT	CAG
		v	1	L	ĸ	R	Ł	Ţ	E	K	D	G	K	V	K	٧>
		630			64	.0		$\epsilon$	50			660			67	0
	•	•				•	•		. •		•	•	,	•		•
	111	1 TG	AAT TTA	GAC	ACT	GCA	GGT	TCT	AAC	AAA	AAA	ACA	GGT	AAA	TGG	GAA
	F	L	N	CIG	TGA T	CGI	CCA	AGA	TIG	TTT						
	•	_	• `	-	•	•	J	3	14	Λ.	K	T	G	K	W	E>
		•	088			690			70	0		7	10			720
	c) c	1.00	•		•			•		•	•		•.		•	•
	CTG	AC1	ACT.	AGC	ACT	TTA	ACA	ATT	AGT	GCT	GAC	AGC	AAA	AAA	ACT	AAA
	D	S	TOA	100	TGA T	YW1	TGI	TAA	TCA	CGA	CIG	TCG	TTT	TTT	TGA	TTT
	_		•	3	•	L	•	1	3	Α.	D	5	K	K	Т	K>
		•	73			. 7	40			750			76	0		
	Cam	4 4	cma	•						•				•	•	
	CLF	TIG.	CIG	ANC	ATT	ACA TCT	GAT	GGT	ACA	TTA	ACA	GTA	CAA	CAA.	TAC	AAC
	CIA	~~~	CAC	$\sim \sim$	WYI	101	CIA	CCA	IGT	TAA	IGT	CAT	GTT	GTT	ATG	TTG
	D	I.	v	F	T.	T	ח	C	T	T	T	37	^	_	Y	

FIGURE 36 (2 of 4)

### B-31 OSP B/ B-31 P41 (140 -295)

770			780			79	0		8	300			810		
•		•	•		•		•	•		•		•	. •		•
				AGC											
TGT	CGA	CCT	TGG	TCG											GAA
T	A	G	T	S	L	E	G	S	A	S	E	I	K	N	L>
82	20		8	30			840			89	0		ε	860	
	◆,	•		•		•	•		•		•	•		•	
				AAC											
AGT	CTC	GAA		TTG								CGA	AGA	GTT	TTA
S	E	L	K	N	A	L	K	G	Ħ,	`P~	H	A	S	Q	N>
	870			88	30		8	90			900		·	91	.0
•	•		•		•	•		•		• .	•		• '	•	•
GTA	AGA	ACA	GCT	GAA	GAG	CTT	GGA	ATG	CAG	CCT	GCA	AAA	TTA	AAC	ACA
CAT	TCT	TGT	CGA	CTT	CTC	GAA	CCT	TAC	GTC	GGA	CGT	TTT	AAT	TTG	TGT
v	R	T	A	E	E	L	G	M	Q	P	A	. K	I	N	T>
	•	920			930			94	10		9	50			960
•	•	•		•	•		•		•	•		•		•	•
CCA	GCA	TCA	CTT	TCA	GGG	CTT	CAA	GCG	TCT	TGG	ACT	ATT	AGA	GTT	CAT
GGT	CGT	AGT	GAA	AGT	CCC	GAA	GTT	CGC	AGA	ACC	TGA	TAA	TCT	CAA	GTA
P	A	s	L	s	G	L	Q	A	S	W	T	L	R	V	H>
		9.	70			80			990			100	00		
	•			•	•	•		•	•		•		•	•	
GTT	GGA	GCA	ACC	CAA	GAT	GAA	GCT	TTA	GCT	GTA	TAA	ATT	TAT	GCA	GCT
				GTT											
V		•		Q			A		A		N	I	Y	A	λ>
1010			1020			10	30		10	040		1	1050		
•		•	•		•		•	•	•	•		•	•		•
AAT	GTT	GCA	TAA	CTT	TTC	TCT	GGT	GAG	GGA	GCT	CAA	ACT	GCT	CAG	GCT
TTA	CAA	CGT	TTA	GAA	AAG	AGA	CCA	CTC	CCT	CGA	GTT	TGA	CGA	GTC	CGY
N	v	A	N	L	F	s	G	E	G	A	Q	T	A	Q	λ>
10	60		1	070			1080			10	90		1:	100	
	•	•		•		•	•		•		•	•		•	
				GAG											
				CTC											
A	P	v	Q	E	G	V	Q	Q	E	G	, A	Q	Q	. P	Α>
	1110	)		11	20		1	130			1140			11	50
•	•		•		•	•		•		•	•		•		•
															GTT
GGA	CGA												CAA	TTA	CAA
P	A	T	Α	P	S	Q	G	G	V	N.	S	P	V	N	٧>

FIGURE 36 (3 of 4)

#### B-31 OSP B/ B-31 P41 (140 -295)

1160 1170 1180 1190 ACA ACT ACA GTT GAT GCT AAT ACA TCA CTT GCT AAA ATT GAA AAT GCT TGT TGA TGT CAA CTA CGA TTA TGT AGT GAA CGA TTT TAA CTT TTA CGA T T T V D A N T S L A K I E N A> 1210 1220 1230 ATT AGA ATG ATA AGT GAT CAA AGG GCA AAT TTA GGT GCT TTC CAA AAT TAA TCT TAC TAT TCA CTA GTT TCC CGT TTA AAT CCA CGA AAG GTT TTA IRMISDQRANLGAFQ N> 1260 1250 1270 1280 AGA CTT GAA TCT ATA AAG AAT AGT ACT GAG TAT GCA ATT GAA AAT CTA TCT GAA CTT AGA TAT TTC TTA TCA TGA CTC ATA CGT TAA CTT TTA GAT R L E S I K N S T E Y A I E N L> 1300 1310 1320 AAA GCA TCT TAT GCT CAA ATA GGT CAC C TTT CGT AGA ATA CGA GTT TAT CCA GTG G K A S Y A Q I G H X>

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87/133

Osp b/ fla (122-234) osp c Sequence Range: 1 to 1765

> . 30 20 10 GCA CAA AAA GGT GCT GAG TCA ATT GGT TCT CAA AAA GAA AAT GAT CTA CGT GTT TTT CCA CGA CTC AGT TAA CCA AGA GTT TTT CTT TTA CTA GAT AQKGAESIGSQKE NDL> 80 90 70 60 • AAC CTT GAA GAC TCT AGT AAA AAA TCA CAT CAA AAC GCT AAA CAA GAC TTG GAA CTT CTG AGA TCA TTT TTT AGT GTA GTT TTG CGA TTT GTT CTG N L E D S S .K K S H Q N A K Q D> 130 120 110 CTT CCT GCG GTG ACA GAA GAC TCA GTG TCT TTG TTT AAT GGT AAT AAA GAA GGA CGC CAC TGT CTT CTG AGT CAC AGA AAC AAA TTA CCA TTA TTT L P A V T E D S V S L F N G N K> 180 170 160 150 ATT TTT GTA AGC AAA GAA AAA AAT AGC TCC GGC AAA TAT GAT TTA AGA TAA AAA CAT TCG TTT CTT TTT TTA TCG AGG CCG TTT ATA CTA AAT TCT I F V S K E K N S S G K Y D L R> 200 210 220 230 GCA ACA ATT GAT CAG GTT GAA CTT AAA GGA ACT TCC GAT AAA AAC AAT CGT TGT TAA CTA GTC CAA CTT GAA TTT CCT TGA AGG CTA TIT TTG TTA A T I D Q V E L K G T · S D E N N> 285 270 260 250 GGT TCT GGA ACC CTT GAA GGT TCA AAG CCT GAC AAG AGT ANA GTA AAA CCA AGA CCT TGG GAA CTT CCA AGT TTC GGA CTG TTC TCA TIT CAT TIT G S G T L E G S K P D K S X V K> 320 310 300 290 • TTA ACA GTT TCT GCT GAT TTA AAC ACA GTA ACC TTA GAA GCA TTT GAT AAT TGT CAA AGA CGA CTA AAT TTG TGT CAT TGG AAT CTT CGT AAA CTA L T V S A D L N T V T L E A F D> 370 360 350 * * * GCC AGC AAC CAA AAA ATT TCA AGT AAA GTT ACT AAA AAA CAG GGG TCA CGG TCG TTG GTT TTT TAA AGT TCA TTT CAA TGA TTT TTT GTC CCC AGT A S N Q K I S S K V T K K Q G S>

> > FIGURE 37 (1 of 5)

Osp b/ fla (122-234) osp c

410 400 420 ATA ACA GAG GAA ACT CTC AAA GCT AAT AAA TTA GAC TCA AAG AAA TTA TAT TGT CTC CTT TGA GAG TTT CGA TTA TTT AAT CTG AGT TTC TTT AAT I T E E T L K A N K L D S K K L> 460 440 450 470 ACA AGA TCA AAC GGA ACT ACA CTT GAA TAC TCA CAA ATA ACA GAT GCT TGT TCT AGT TTG CCT TGA TGT GAA CTT ATG AGT GTT TAT TGT CTA CGA TRSNGTTLEYSQITD A> , a .. = 15 1 ... 500 510 GAC AAT GCT ACA AAA GCA GTA GAA ACT CTA AAA AAT AGC ATT AAG CTT CTG TTA CGA TGT TTT CGT CAT CTT TGA GAT TTT TTA TCG TAA TTC GAA D N A T K A V E T L K N S I K L> 540 530 550 560 570 GAA GGA AGT CTT GTA GTC GGA AAA ACA ACA GTG GAA ATT AAA GAA GGT CTT CCT TCA GAA CAT CAG CCT TTT TGT TGT CAC CTT TAA TTT CTT CCA EGSLVVGKTTVEIKEG> 580 590 600 610 ACT GTT ACT CTA AAA AGA GAA ATT GAA AAA GAT GGA AAA GTA AAA GTC TGA CAA TGA GAT TTT TCT CTT TAA CTT TTT CTA CCT TTT CAT TTT CAG T V T L K R E I E K D G K V K V> 640 650 630 660 • TTT TTG AAT GAC ACT GCA GGT TCT AAC AAA AAA ACA GGT LAA TGG GAA AAA AAC TTA CTG TGA CGT CCA AGA TTG TTT TTT TGT CCA TTT ACC CTT F L N D T A G S N K K T G K W E> 680 690 710 GAC AGT ACT AGC ACT TTA ACA ATT AGT GCT GAC AGC AAA AAA ACT AAA CTG TCA TGA TCG TGA AAT TGT TAA TCA CGA CTG TCG TTT TTT TGA TTT D S T S T L T I S A D S K K T K> 740 760 . GAT TTG GTG TTC TTA ACA GAT GGT ACA ATT ACA GTA CAA CAA TAC AAC CTA AAC CAC AAG AAT TGT CTA CCA TGT TAA TGT CAT GTT ATG TTG 

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89/133

Osp b/ fla (122-234) osp c

790 800 810 780 770 • • ACA GCT GGA ACC AGC CTA GAA GGA TCA GCA AGT GAA ATT AAA AAT CTT TGT CGA CCT TGG TCG GAT CTT CCT AGT CGT TCA CTT TAA TTT TTA GAA T A G T S L E G S A S E I K N L> 850 820 830 840 TCA GAG CTT AAA AAC GCT TTA AAA GGT CAC CCC ATG GGA AAT AAT TCA AGT CTC GAA TTT TTG CGA AAT TTT CCA GTG GGG TAC CCT TTA TTA AGT S E L K N A L K G H P-M G K N S> 880 890 900 870 • • GGG AAA GAT GGG AAT ACA TCT GCA AAT TCT GCT GAT GAG TCT GTT AAA CCC TTT CTA CCC TTA TGT AGA CGT TTA AGA CGA CTA CTC AGA CAA TTT G K D G N T S A N S A D E S V K> 950 930 940 920 • GGG CCT AAT CTT ACA GAA ATA AGT AAA AAA ATT ACG GAT TCT AAT GCG CCC GGA TTA GAA TGT CTT TAT TCA TTT TTT TAA TGC CTA AGA TTA CGC G P N L T E I S K K I T D S N A> 980 990 1000 970 GTT TTA CTT GCT GTG AAA GAG GTT GAA GCG TTG CTG TCA TCT ATA GAT CAA AAT GAA CGA CAC TTT CTC CAA CTT CGC AAC GAC AGT AGA TAT CTA V L L A V K E V E A L L S S I D> 1020 1030 1040 GAA ATT GCT GCT AAA GCT ATT GGT AAA AAA ATA CAC CAA AAT AAT GGT CTT TAA CGA CGA TTT CGA TAA CCA TTT TTT TAT GTG GTT TTA TTA CCA EIAAKAIGKKIHQNNG> 1070 1080 1090 1090 TTG GAT ACC GAA TAT AAT CAC AAT GGA TCA TTG TTA GCG GGA CGT TAT AAC CTA TGG CTT ATA TTA GTG TTA CCT AGT AAC AAT CGC CCT GCA ATA L D T E Y N H N G S L L A G R Y> 1120 1130 1140 1110 GCA ATA TCA ACC CTA ATA AAA CAA AAA TTA GAT GGA TTG AAA AAT GAA CGT TAT AGT TGG GAT TAT TTT GTT TTT AAT CTA CCT AAC TTT TTA CTT A I S T L I K Q K L D G L K N E>

FIGURE 37 (3 of 5)

Osp b/ fla (122-234) osp c 1170 1280 1190 1200 1160 GGA TTA AAG GAA AAA ATT GAT GCG GCT AAG AAA TGT TCT GAA ACA TTT CCT AAT TTC CTT TTT TAA CTA CGC CGA TTC TTT ACA AGA CTT TGT AAA G L K E K I D A A K K C S E T F> 1210 1220 1230 1240 ACT AAT AAA TTA AAA GAA AAA CAC ACA GAT CTT GGT AAA GAA GGT GTT TGA TTA TTT AAT TTT CTT TTT GTG TGT CTA-GAA-CCA TTT CTT CCA CAA T N K L K E K H T D L G K E G V> 1260 1270 1280 1290 1250 ACT GAT GCT GAT GCA AAA GAA GCC ATT TTA AAA ACA AAT GGT ACT AAA TGA CTA CGA CTA CGT TTT CTT CGG TAA AAT TTT TGT TTA CCA TGA TTT T D A D A K E A I L K T N G T K> 1310 1320 1330 ACT AAA GGT GCT GAA GAA CTT GGA AAA TTA TTT GAA TCA GTA GAG GTC TGA TTT CCA CGA CTT CTT GAA CCT TTT AAT AAA CTT AGT CAT CTC CAG T K G A E E L G K L F E S V E V> 1360 1370 1380 TTG TCA AAA GCA GCT AAA GAG ATG CTT GCT AAT TCA GTT AAA GAG CTT AAC AGT TTT CGT CGA TTT CTC TAC GAA CGA TTA AGT CAA TTT CTC GAA L S K A A K E M L A N S V K E L> 1420 1400 1410 1430 ACA AGC CCT GTT GTG GCA GAA AGT CCA AAA AAA CCT GGT ACC ATG GCT TGT TCG GGA CAA CAC CGT CTT TCA GGT TTT TTT GGA CCA TGG TAC CGA T S P V V A E S P K K P G T M A>

1450 1460 1470 1480

CAA TAT AAC CAA ATG CAC ATG TTA TCA AAC AAA TCT GCT TCT CAA AAT GTT ATA TTG GTT TAC GTG TAC AAT AGT TTG TTT AGA CGA AGA GTT TTA Q Y N Q M H M L S N K S A S Q N>

1490 1500 1510 1520 1530

GTA AGA ACA GCT GAA GAG CTT GGA ATG CAG CCT GCA AAA ATT AAC ACA CAT TCT TGT CGA CTT CTC GAA CCT TAC GTC GGA CGT TTT TAA TTG TGT V R T A E E L G M Q P A K I N T>

1.1

### 91/133

Osp b/ fla (122-234) osp c

1550 . 1560 1570 CCA GCA TCA CTT TCA GGG CTT CAA GCG TCT TGG ACT TTA AGA GTT CAT GGT CGT AGT GAÁ AGT CCC GAA GTT CGC AGA ACC TGA AAT TCT CAA GTA PASLSGLQASWTLRVH> 1600 1610 1620 1630 1590 GTT GGA GCA ACC CAA GAT GAA GCT ATT GCT GTA AAT ATT TAT GCA GCT CAA CCT CGT TGG GTT CTA CTT CGA TAA CGA CAT TTA TAA ATA CGT CGA V G A T Q D E A I A V N I Y A A> 1650 1660 1670 1680 1640 AAT GTT GCA AAT CTT TTC TCT GGT GAG GGA GCT CAA ACT GCT CAG GCT TTA CAA CGT TTA GAA AAG AGA CCA CTC CCT CGA GTT TGA CGA GTC CGA N V A N L F S G E G A Q T A Q A> 1690 1700 1710 1720 GCA CCG GTT CAA GAG GGT GTT CAA CAG GAA GGA GCT CAA CAG CCA GCA

CCT GCT ACA GCA CCT TCT CAA GGC GGA GTT GGT CAC C
GGA CGA TGT CGT GGA AGA GTT CCG CCT CAA CCA GTG G
P A T A P S Q G G V G H X>

				10			20			30				40		
OspC-B31	ATG TAC	AAA TTT	AAG	AAT	ACA	TTA	AGT	GCG	ATA	TTA	ATG	ACT	TTA		* ATT AAT	TTT AAA
1. OspC-P1 [ 1832 ]	κ	•••	•••	LO •••	•••	•••	20	•••	•••	30	•••	•••	•••	40	•••	>
2. OspC-TI [ 1786 ]																
3. OspC-K4 [ 1774 ]	• • • •	•••	•••		•••	•••	20	•••	•••	30	•••	•••	•••	40	•••	>
	50		_	60				70	•.		80		_	90		
OspC-831	ATA	TCT	TGT	AAT	AAT	TCA	GGG	AAA	GAT	GGG	AAT	ACA TGT	TCT	GCA CGT	AAT	TCT
1. OspC-PF [ 1832 ]	(50	•••	•••	60 •g•	•••	•••	•	70	.g.				9Ó	a.t	•••	c>
2. OspC-TF [ 1786 ]	R50			60	• • •		t	70 999			tc.,	80 g	•••	a.t		0>
3. OspC-K4																
•	10	00		1	110			120			13	30		1	.40	
OspC-B31	GCT	GAT	GAG	TCT	GTT	AAA	GGG	CCT	TAA	CTT	ACA	GAA	ATA	AGT TCA	AAA	AAA TTT
1. OspC-10	00	с		10	.ca		120	•••		13	0			40		>
2. OspC-TF [ 1786. ]																
3. OspC-K4																
OspC-B31														* GTT		*
																000
1 Cancant		TGC	CTA		ATT	CGC	CAA	AAT	GAA	CGA	CAC	TTT	CTC		CTT	CGC
1. OspC-PF [ 1832 ]	(150	TGC	CTA	16	TTA 50	CGC	CAA	AAT 170 9	•••	CGA	180 t	TTT	a	19	CTT 0 •••g	a.t>
1. OspC-PF [ 1832 ] 2. OspC-T] [ 1786 ]	(150	TGC	CTA	16	TTA 50	CGC	CAA	AAT 170 9	•••	CGA	180 t	TTT	a	19	CTT 0 •••g	a.t>
[ 1001 ]	140	TGC a	CTA	150 150	TTA 50	a	t	AAT 170 9 50	 g	CGA	180 t .70		a	19  180 	CTT 0 ··g	a.t>
2. OspC-T] [ 1786 ] 3. OspC-K]	(150  140 	··a ··a	CTA	150  150	TTA 50 	a a	t	AAT 170 9 50 	g	CGA	180 t .70		a	19 180  180	CTT O O O O G O G O G O G O G O G O G O G	a.t>t>
2. OspC-T] [ 1786 ] 3. OspC-K]	(150  140  140 	TGCaa CTG	CTA 200 *TCA	160  150 	TTA 50	aa 210 GAT	t t t	AAT 170 9 50  9 ATT	g g g	CGA	70 	TTT	a a a	19  180 	CTT 0ggg	a.t>t>t> 240
2. OspC-TI [ 1786 ] 3. OspC-KI [ 1774 ]	(150  140  140 	TGCaa CTG	CTA 200 TCA AGT	150  150 	TTA 50 ATA TAT	aa 210 GAT CTA	t 16 t GAA CTT	AAT 170 9 60 60 g ATT TAA	g g g g	CGA   1  20  GCT CGA	70  .70 	TTT 2 GCT CGA	aaaaaaa	19  180  180 	CTT O ··g ··g ··g AAA TTT	a.t>t>t> 240 AAA TTT

Figure 38 (1 of 3)

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93/133

3. OspC-K	(4 19	0		2	00			210			22	0		2	30	at >
[ 1774 ]	•••	a.c	• • •	• • •	• • •	• • •	• • •	C	• • •	aa.	• • •	•••	•••	•••	• • •	90.,
		*	25	0	*	2	60 . *		*	270		*	28	*	*	
OspC-B31	TAT	GTG	CAA .	AAT ATT	AAT TTA	GGT CCA	TTG AAC	GAT CTA	ACC TGG	GAA CTT	TAT ATA	AAT ATT	CAC GTG			
1. OspC-F	PK .	25	0		. 2	60			270 g.t	tt.	a	28	30 - • ¤		2	90
[ 1832 ]	•••	g	a.t	•••		tac	•••	•••	9.0							
				240			0		•	260			270			•
2. OspC-1	rr •••			240	g		a	•••	.a.	•••	gca	• • •	.ga	c	.a.	>
3. OspC-F		240			25	n		- 2	260			270			28	10
	290			300			31	LO		:	320		-	330	-	
OspC-B31		~~~	~~~	~~~	COT	ጥልጥ	CCD	ATA	TCA	ACC	CTA	ATA	AAA	CAA	AAA	TTA
Ospc-ssr	AAC	AAT	CGC	CCI	GCA	WTW.	CG 1							GTT	TTT	TAA
1. OspC-1 [ 1832 ]	PK	• • •	a	• • •	gcc	• • •	• • •	• • •	• • •	• • •	•••		•••	g	•••	g>
2. OspC-3	280	a		90	ac.		300 .a.			3. .aa	10	• • •	.c.	320	•••	>
3 Oct.C=.	 КЛ		90	•••	30.	300			3	10		:	320			330
3. OspC-1	•••	•••	a	• • •	gcc	•••	•••	•••	• • •	•••	•••	•••	.c.	g	•••	>
	. 3	40		3	350			360			31	70	*	3	880	
OspC-B31	GAT CTA		mma		B B W	CDD	CCA	ጥጥል	DAG	GAA	AAA	ATT	GAT	GCG	GCT	AAG TTC
					ttt						•					
1. OspC- [ 1832 ]	340	22		350	\ a	360	.a.		3	70 ac.	g		380 .ca	aa.		390 >
2 00-	mn 2 2 A			3.	40			350			360			31	70	
[ 1786 ]	ag.	.t.	• • •	ŧ	tca	•••	.a.	•••	•••	a	•••	•••	a.a	.a.	•••	>
					ttc										•	
3. OspC-	·K4	aa.	3	40	. a	350	.ag	• • • •	360	) : a			70 a	.a.	•••	380
( 2.77	3 -															
	*	390 *		*	4	00	*		410		*	420		*		30 *
OspC-B31	AAA TTT	TGT ACA	TCT	GAA	ACA TGT	AAA	ACI TGA	TTA	TTT	ATT A	AAA Z	GAA	TTT	GTG	TGT	GAT
1. OspC- [ 1832 ]	-PK		4	00	a s		410			420	)	agt		30 t	g	>
2 0000	_TD	200			390	1		4	100			410			420	)
[ 1786 ]	g.t			:	.a.	•••	•••	.c.	(	g c.	• • • •	t	.gt	t	g	g>
3. OspC- [ 1774 ]	-K4	c ca.	390	) 	g		100		g	410 . c.		.gt	420 tct	t	g	c.a>
			440			450	)			460			470			480

Figure 38 (2 of 3)

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OspC-B31	GAA	CCA	TTT	CTT	CCA	CAA	TGA	CTA	CGA	CTA	CGT	TTT	CII	CGG	TAA	AAT
1. OspC-F [ 1832 ]	440	c		450 c.g	.a.	.c.	c	50	.a.	c	70	•••	.c.	480 t	•••	>
2. OspC-1 [ 1786 ]																
3. OspC-4 [ 1774 ]	30	a	gtt	440 .ct	.c.	.c.	450	•••	.a.	c		•••	•••	170 t	•••	>
		•	49	90	•	9	500		*	510		*	52	20	*	• .
OspC-B31	TTT	ACA TGT	AAT TTA	GGT CCA	ACT TGA	AAA TTT	ACT TGA	AAA TTT	CCA	CGA	GAA CTT	GAA CTT	CTT GAA	GGA CCT	AAA TTT	TTA AAT
1. OspC-P [ 1832 ]	K 49	90	c	.ca	500	.cc	ga.	510	•••	•••	52 a	20	t	aa.	30 g.t	>
2. OspC-1 [ 1786 ]	R.		480 c	a		49	90 gac	g		500	a		510	.a.	g.g	>
3. OspC-K [ 1774 ]	4480 ••g	t		49 cc.	90	•••	ga.	500 g	•••	•••	510 a	.c.	•••	52 aa.	g.c	>
	530 *		*	540		*	5	50	*	5	60		* i	570		*
OspC-B31	TTT AAA	GAA CTT	TCA AGT	GTA CAT	GAG CTC	GTC CAG	TTG AAC	TCA AGT	AAA TTT	GCA CGT	GCT CGA	AAA TTT	GAG CTC	ATG TAC	CTT GAA	GCT CGA
1. OspC-F [ 1832 ]																
2. OspC-5 [ 1786 ]	20	a	!	530 c	a	ag.	540		•••	55	g	с	.ca	660 gca	t.a	a>
														-		
3. OspC-K																
	(4 .c.	530	•••	•••	540 a	ag.	•••	5: g.,	50	•••	g	60 c	a	gca	570 t.a	
	.c. 51	530  80 *	**		540 a 590 *	ag.	*	5: g., 600 *	SO	* GTT	g 61	660 C LO *	a *	gca GAGT	570 t.a 520	> AAA
3. OspC-R [ 1774 ] OspC-B31	C4 .C. SI AAT TTA	30 30 * TCA AGT	* GTT CAA	AAA TTT	540 a 590 * GAG CTC	ag. CTT GAA	* ACA	g 600 * AGC TCG	CCT GGA	* GTT CAA	GTG CAC	660 C LO * GCA CGT	a * GAA CTT	gca AGT TCA	570 t.a 520 * CCA GGT	AAA TTT
3. OspC-F [ 1774 ] OspC-B31 1. OspC-F [ 1832 ]	SAAT TTA	BO TCA AGT	GTT CAA 590	AAA TTT	540 a 590 * GAG CTC	ag. CTT GAA 600	* ACA TGT	g 600 * AGC TCG	CCT GGA 6:	GTT CAA	61 GTG CAC	GCA CGT	GAA CTT	gca AGT TCA	570 t.a 520 * CCA GGT	AAA TTT 630
3. OspC-K [ 1774 ]  OspC-B31  1. OspC-F [ 1832 ]  2. OspC-7 [ 1786 ]  3. OspC-F	AAT TTA PK	30 * TCA AGT	GTT CAA 590	AAA TTT	540 a 590 * GAG CTC	ag. CTT GAA 600	* ACA TGT	9 600 *AGC TCG t	CCT GGA 6:	GTT CAA	61 GTG CAC	GCA CGT	GAA CTT	gca AGT TCA	570 t.a 520 * CCA GGT	AAA TTT 630
3. OspC-K [ 1774 ]  OspC-B31  1. OspC-F [ 1832 ]  2. OspC-7 [ 1786 ]	AAT TTA PK	30 * TCA AGT	GTT CAA 590	AAA TTT	540 a 590 GAG CTC a	ag. CTT GAA 600	* ACA TGT	9 600 *AGC TCG t	CCT GGA 6:	GTT CAA	61 GTG CAC	GCA CGT	GAA CTT	gca AGT TCA	570 t.a 520 * CCA GGT	AAA TTT 630
3. OspC-K [ 1774 ]  OspC-B31  1. OspC-F [ 1832 ]  2. OspC-7 [ 1786 ]  3. OspC-F [ 1774 ]	50 AAT TTA PK CR570 C4	30 * TCA AGT	GTT CAA 590	AAA TTT	540 a 590 * GAG CTC	ag. CTT GAA 600	* ACA TGT	9 600 *AGC TCG t	CCT GGA 6:	GTT CAA	61 GTG CAC	GCA CGT	GAA CTT	gca AGT TCA	570 t.a 520 * CCA GGT	AAA TTT 630
3. OspC-F [ 1774 ]  OspC-B31  1. OspC-F [ 1832 ]  2. OspC-F [ 1786 ]  3. OspC-F [ 1774 ]	AAT TTA PK CR570 C4 AAA TTT	30 * TCA AGT	GTT CAA 590	AAA TTT	540 a 590 * GAG CTC a	ag. CTT GAA 600	* ACA TGT	9 600 *AGC TCG t	CCT GGA 6:	GTT CAA	61 GTG CAC	GCA CGT	GAA CTT	gca AGT TCA	570 t.a 520 * CCA GGT	AAA TTT 630
3. OspC-K [ 1774 ]  OspC-B31  1. OspC-F [ 1832 ]  2. OspC-7 [ 1786 ]  3. OspC-F [ 1774 ]	AAT TTA PK CR570 C4 AAA TTT	TCA AGT	GTT CAA 590	AAA TTT	540 a 590 * GAG CTC a	ag. CTT GAA 600	* ACA TGT	9 600 *AGC TCG t	CCT GGA 6:	GTT CAA	61 GTG CAC	GCA CGT	GAA CTT	gca AGT TCA	570 t.a 520 * CCA GGT	AAA TTT 630
3. OspC-F [ 1774 ]  OspC-B31  1. OspC-F [ 1832 ]  2. OspC-F [ 1786 ]  3. OspC-F [ 1774 ]  OspC-B31  1. OspC-F	AAT TTA PK AAA TTT	30 * TCA AGT 	GTT CAA 590  80 	AAA TTT	540 a 590 * GAG CTC a	ag. CTT GAA 600	* ACA TGT	9 600 *AGC TCG t	CCT GGA 6:	GTT CAA	61 GTG CAC	GCA CGT	GAA CTT	gca AGT TCA	570 t.a 520 * CCA GGT	AAA TTT 630

Figure 38 (3 of 3)
SUBSTITUTE SHEET (RULE 26)

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BO ospD	CTA C	TG I						mma		ጥሮሽ	מידים	di Ciri	4.(2.1.	TCT	TTA DDT	CTA
1. P-Gau				_			20			30			4	0		
1. P-Gau	o		1	U						• • •	• • •	• • •	• • •	.a.	• • •	•••>
[ 2804 ]	•••	• • •	• • •	• • •	• • •	• • •							4	^		
2. DK29 o [ 2786 ]	8		1	0			20			30			~ 4	U		>
r 2786 ]	•••	0	z	• • •	•••	• • •	• • •	• • •	•••	•••	•••	•••	9	•••	•••	
3. K48 OB				^			20			30			4	0		
3. K48 OB	р						•••	• • •		• • •	•••	• • •	• • •	• • •	•••	>
[ 2/86 ]	•••	•••	• • •	•••	-											
								70		,	80			90		
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BO ospD																
Bo osbn	AAT ATT	CTT (	CCA	CAT	TTG	AGT	TTT	CTA	ATG	CTC	AGT	TTT	TTT.	GTC	TCA	TAT
											80			90		
1. P-Gau [ 2804 ]	o50			60				70						•••		>
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2. DK29 c [ 2786 ]	-E0			60				70			80	•		90		_
1 2786 1	3830								• • •	• • •	• • •	• • •	• • •	• • •	•••	>
( 2700 )	•••							~^			20			90		
3. K48 os	3p50			60				70								>
[ 2786 ]	• • •	• • •	• • •	• • •	٠g٠	• • •	• • •	• • •	•••							
	10	00			110			120	)	_	1	30	*	•	140	
BO ospD								_ mm/		ת מים י	דים מ	מיז מי	ייאמ	TICA	CIA	AAA
BO ospD	CTA GAT	GGT	GAA	TTA	AA'I	CAC	CIA	r AAC	CCC	GTI	TGA	TGT	TTA	AGT	GAT	TTT
	GAT	CCA	CTT	AMI	114	. 01.										
1. P-Gau [ 2804 ]	0 10	00			110			120	)		1	.30			140	>
[ 2804 ]	•••				• • •	• • • •		• • • •	• • • •	•••	•••	• • •	• • •	•••	• • •	
								120	<b>`</b>		1	30			140	
2. DK29 { 2786 }	os 10	00			110				• •••						• • •	>
[ 2/80 ]	• • •	• • •	•••	•••							_				3.40	
3. K48 o	sp 1	00			110			12	0		]	130			140	>
[ 2786 ]	••••	• • •	• • •	• • •	• • •	• ••		• ••	• • •	• • • •	• • • •	• • • •	•••	• • •	•••	• • •
		150				160			170			180	)		1	.90
	*	*		*		*		*	<b>*</b>		*		t 	· GAG	GCA	TAA
BO ospD	GAA	GCA	AAA	AA'	r ac	A AC	A GA	AA T.	T TT	ጸ ጸጸ ጥ ጥጥ	T GC	T AG	TTP	CTC	CGI	TTA
	CTT	CGT	TTI	TT	A TG	T TG	T CI	H II	A AA	1 11	n co.	1 110.				
1 D-Car		150	1			160			170			180	0			190
1. P-Gav [ 2804 ]		150										• • •	• • • •	• • •	•••	>
( 2004 )		• • • •	-									10	^			190
2. DK29	OS	150	)			160			170	'		10				>
2. DK29 [ 2786 ]	• • •	•••	• •	• • •	• ••		• • •	• • •								
3. K48	222	150	,			160			170	)		18	0			190
3. K48 ( [ 2786 ]	neħ													• • • •	• ••	>
( =, = ,																
			000			<b>3</b> .	10			220			230			240
		<b>k</b>	200		*	2.	*	,	k	*		*	. *			*
BO ospD			r GT	A G		~ ~	TOTAL BY	מ מיד	TT G	ന്ദ്ര ദീ	א די	TT TI	'A AT	T TC	A TC	T GCT A CGA
no oahn	TT	T CA	A CA	T C	T C	GT C	AA T	AT T	CA C	AC C	AA TI	AA A?	AT TA	A AG	T AG	A CGA
	_									•						

2. DK29 os

390

										96	//3:	?					
	1. P-Gau	0	•	200			210							230			240
Į	1. P-Gau 2804 ]	•••	•••	•••	• • •	• • •	•••	• • •	• • •	• • •	• • •	• • •	• • •	• • •	• • •	• • •	>
	2. DK29 c	В	2	200			210			2:	20		:	230			240
ſ	2786 ]	• • •	•••	•••	• • •	• • •	•••	• • •	•••	• • •	• • •	• • •	• • •	• • •	• • •	• • •	>
	3. K48 os	3p	2	200			210			2:				230			240
Į	2786 ]	• • •	•••	•••	• • •	• • •	• • •	• • •	• • •	•••	• • •	•••	• • •	•••	•••	• • •	>
			*	25	50 ★	*	- 1	260 *		*	270 *		*	28	30 *	*	
B	O ospD	GCA CGT	GAT CTA	CAG GTC	GTA CAT	AAA TTT	GGT CCA	CAA GTT	CAA GTT	CAA GTT	ATA TAT	TGC ACG	ACG TGC	ATT TAA	TAG ATC	CTC GAG	AAA TTT
	1. P-Gau	0		25	50		:	260			270			28	30		
I,	2804 ]	•••	• • •	• • •	• • •	• • •	• • •	• • •	• • •	• • •	• • •	• • •	• • •	•••	• • •	• • •	>
	2. DK29 c	8		25	50		2	260			270			28	30		
ſ	2786 ]	• • •	•••	• • •	•• <b>g</b>	•••	• • •	• • •	• • •	• • •	• • •	• • •	•••	• • •	•••	• • •	>
	3. K48 os	p		29	50		:	260			270			28			
ĺ	2786 J	• • •	• • •	• • •	•• <b>g</b>	• • •	• • •	• • •	• • •	• • •	• • •	• • •	•••	• • •	•••	• • •	>
		290 *													330		*
B	OgpO	TGG ACC	CAG GTC	AAA TTT	TAG ATC	ATT TAA	TAG ATC	AAA TTT	AAA TTT	TAA ATT	AGG TCC	AAT	CTA GAT	GTG CAC	ATA TAT	AAG TTC	TAA ATT
	1. P-Gau	290			200			3.	10			320			330		
	2804 ]														_	• • •	>
	2. DK29 c	290			300			3:	10		:	320			330		
ι	2786 j	•••	• • •		•••			• • •	•••	• • •	• • • •	•••	•••	••,•		• • •	>
	3. K48 os	290			300			3:	10		:	320			330		
E	2786 ]	• • •		• • •											•••	•••	>
															_		
		34	40 *	*		350 *		*	360 *		*	37	70 *	*	3	180 *	
В	Ogao C	TAG	TTG	CGG	CTA	ATG	TTG	CGA	AAG	AAG	CAT	ATA	ACC	TTA	CTA	AAG	CAG
		ATC	AAC	GCC	GAT	TAC	AAC	GCI	110	110	GIA	IMI	166	AAI	GAI	110	GIC
r	1. P-Gau 2804 ]	0 34	40		:	350			360			3.	70			80	>
·				•••	•••	•••	•••	•••	•••	•••	•••			•••			
ſ	2. DK29 c	ов 34	40			350			360			31	70			380	>
•	•	•	40			250			260			٠.			_	200	
ĺ	3. K48 of 2786 ]	•••	•••		•••	350	•••	•••	360	•••	•••		70	•••	•••	380	>
			200				00			440			400				20
		*	390 *		*	-	00 *	*	•			*	420 *		*		30 *
B	O ospD		AAC TTG														
	1. P-Gau						00			410			420				30
(	2804 ]	•••		•••	•••	• • •	•••	•••	• • •	•••	• • •	•••	•••	• • •	• • •	•••	>

Figure 39 (2 of 4)
SUBSTITUTE SHEET (RULE 26)

400

410

420

430

`				97/1	33		
[ 2786 ]			•••	•••	•••		
3. K48 osp	390	4	00	410		420	430
[ 2700 ]	•••	•••					
	440		450	4	60	470 *	480
BO OBPD AAC	ACT ATC	TGA TTC	TGA TGA	AAC AGA	ACG AGT	TTC TGA TG	AAA TAA A
TTG						AAG ACT AC	TTA TTT
1. P-Gau o [ 2804 ]	440		450	4	60	470	480 • • • • • • • • • • • • • • • • • • •
-							
2. DK29 os [ 2786 ]		•••	•••	.g	••••	••• •••	>
3. K48 osp	440		450	4	60	470	480
[ 2786 ]	•••	•••	•••	.g	•••.	••• ••• •••	• •••
	49	90	500		510	520 * *	
BO ospD ACA	* AGC TAA	AGA GGC	TGT AGA	AAT AGC	TTG GAA	AGC CAC AG	T AAA AGT
TGT	TCG ATT	TCT CCG	ACA TC	TTA TCG	AAC CTT	TCG GTG TC	A TTT TCA
1. P-Gau o [ 2804 ]					510	520	>
2. DK29 os [ 2786 ]	49	90	500		510	520	>
3. K48 osp	4	90	500		510	520	
[ 2786 ]	•••	• • • • • • • • • • • • • • • • • • • •	• • • • • •	•••	•••	•••	>
530	*	540	9	550	560	570	)
* BO ospD AAA	* AGA TGA	* GTT AAT	* TGA TG	* * AAA	TGC AGT	CAA AGA GG	* * CATT GGA
						GTT TCT CC	
1. P-Gau 530 [ 2804 ]		540	9	550	560	570	
2. DK29 o530 [ 2786 ]		540	•••	550	560	570	>
3. K48 os530		540	9	550	560	570	0
[ 2786 ]	•••	•••	•••		•••	•••	>
5		590 *		600	6:	10	620
	AAT AAA	GAC AGA		GAA CAA	TAC AAA	ACT TAC AG	
			TTG GC			TGA ATG TC	
1. P-Gau o 5 [ 2804 ]		590 • • • • • •	•••		6:		>
2. DK29 os 5 [ 2786 ]		590	•••	600	6:	10	620
3. К48 овр 5							

									9	8/1	33					
	•	630			64	40			550	•		660		•	61	70
	*	*		*					*			*				
BO ospD	AGA TCT	AGT TCA	AGC TCG	AGA TCT	GTT CAA	AGT TCA	TAA	ACA TGT	GAT CTA	AGC TCG	CAA GTT	AAA TTT	TGT ACA	AGC TCG	GGA CCT	TTA TTA
1. P-Gau	0	630			64	40		(	550			660			6:	70
[ 2804 ]	•••	• • •	• • •	• • •	• • •	• • •	•••	• • •	a		•••					>
2. DK29 o	8	630			64	10		(	550			660			61	70
[ 2786 ]	• • •	• • •	• • •	• • •	•••	• • •	• • •	•••	a	• • •	•••	•••	• • •	•••	•••	>
3. K48 os	p	630			64	10		(	550			660				70 ·
[ 2786 ]	•••	• • •	• • •	•••	• • •	•••	•••	• • •	a	•••	• • •	•••	•••	• • •	•••	>
										`						
	*	•	580		_	690		*	70	<b>*</b>						
BO ospD	AGC	GCA CGT	AGA	AGT TCA	TGT	GGC	CTT GAA	GTT	AAA	TAC	TT AA			٠ ٠.		
1. P-Gau	0		580			690			70	00						
[ 2804 ]											>	•				
2. DK29 o [ 2786 ]	s •••	•••	580			690 •••			70		>					
3. K48 os [ 2786 ]	p	(	680			690			70							

40

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99/133

P41

Sequence Range: 1 to 1011

30 20 ATG ATT ATC AAT CAT AAT ACA TCA GCT ATT AAT GCT TCA AGA AAT AAT

TAC TAA TAG TTA GTA TTA TGT AGT CGA TAA TTA CGA AGT TCT TTA TTA Met Ile Ile Asn His Asn Thr Ser Ala Ile Asn Ala Ser Arg Asn Asn>

90 70 80 60

GGC ATT AAC GCT GCT AAT CTT AGT AAA ACT CAA GAA AAG CTT TCT AGT CCG TAA TTG CGA CGA TTA GAA TCA TTT TGA GTT CTT TTC GAA AGA TCA Gly Ile Asn Ala Ala Asn Leu Ser Lys Thr Gln Glu Lys Leu Ser Ser>

130 120 110 •

GGC TAC AGA ATT AAT CGA GCT TCT GAT GAT GCT GCT GGC ATG GGA GTT CCG ATG TCT TAA TTA GCT CGA AGA CTA CTA CGA CGA CCG TAC CCT CAA Gly Tyr Arg Ile Asn Arg Ala Ser Asp Asp Ala Ala Gly Met Gly Val>

170 180 160 150 •

TCT GGT AAG ATT AAT GCT CAA ATA AGA GGT TTG TCA CAA GCT TCT AGA AGA CCA TTC TAA TTA CGA GTT TAT TCT CCA AAC AGT GTT CGA AGA TCT Ser Gly Lys Ile Asn Ala Gln Ile Arg Gly Leu Ser Gln Ala Ser Arg>

230 220 210 200

AAT ACT TCA AAG GCT ATT AAT TTT ATT CAG ACA ACA GAA GGG AAT TTA TTA TGA AGT TTC CGA TAA TTA AAA TAA GTC TGT TGT CTT CCC TTA AAT Asn Thr Ser Lys Ala Ile Asn Phe Ile Gln Thr Thr Glu Gly Asn Leu>

> 260 270

AAT GAA GTA GAA AAA GTC TTA GTA AGA ATG AAG GAA TTG GCA GTT CAA TTA CTT CAT CTT TTT CAG AAT CAT TCT TAC TTC CTT AAC CGT CAA GTT Asn Glu Val Glu Lys Val Leu Val Arg Met Lys Glu Leu Ala Val Gln>

330 320 310 300 290

TCA GGT AAC GGC ACA TAT TCA GAT GCA GAC AGA GGT TCT ATA CAA ATT AGT CCA TTG CCG TGT ATA AGT CTA CGT CTG TCT CCA AGA TAT GTT TAA Ser Gly Asn Gly Thr Tyr Ser Asp Ala Asp Arg Gly Ser Ile Gln Ile>

370 360

GAA ATA GAG CAA CTT ACA GAC GAA ATT AAT AGA ATT GCT GAT CAA GCT CTT TAT CTC GTT GAA TGT CTG CTT TAA TTA TCT TAA CGA CTA GTT CGA Glu Ile Glu Gln Leu Thr Asp Glu Ile Asn.Arg Ile Ala Asp Gln Ala> WO 95/12676 PCT/US94/12352

100/133

410 390 400 420 CAA TAT AAC CAA ATG CAC ATG TTA TCA AAC AAA TCT GCT TCT CAA AAT GTT ATA TTG GTT TAC GTG TAC AAT AGT TTG TTT AGA CGA AGA GTT TTA Gln Tyr Asn Gln Met His Met Leu Ser Asn Lys Ser Ala Ser Gln Asn> 440 450 460 470 480 GTA AGA ACA GCT GAA GAG CTT GGA ATG CAG CCT GCA AAA ATT AAC ACA CAT TCT TGT CGA CTT CTC GAA CCT TAC GTC GGA CGT TTT TAA TTG TGT Val Arg Thr Ala Glu Glu Leu Gly Met Gln Pro Ala Lys Ile Asn Thro 490 510 ___. 520 CCA GCA TCA CTT TCA GGG CTT CAA GCG TCT TGG ACT TTA AGA GTT CAT GGT CGT AGT GAA AGT CCC GAA GTT CGC AGA ACC TGA AAT TCT CAA GTA Pro Ala Ser Leu Ser Gly Leu Gin Ala Ser Trp Thr Leu Arg Val His> 530 540 550 560 GTT GGA GCA ACC CAA GAT GAA GCT ATT GCT GTA AAT ATT TAT GCA GCT CAA CCT CGT TGG GTT CTA CTT CGA TAA CGA CAT TTA TAA ATA CGT CGA Val Gly Ala Thr Glm Asp Glu Ala Ile Ala Val Asn Ile Tyr Ala Ala> 580 590 600 610 AAT GTT GCA AAT CTT TTC TCT GGT GAG GGA GCT CAA ACT GCT CAG GCT TTA CAA CGT TTA GAA AAG AGA CCA CTC CCT CGA GTT TGA CGA GTC CGA Asn Val Ala Asn Leu Phe Ser Gly Glu Gly Ala Gln Thr Ala Gln Ala> 640 650 660 GCA CCG GTT CAA GAG GGT GTT CAA CAG GAA GGA GCT CAA CAG CCA GCA CGT GGC CAA GTT CTC CCA CAA GTT GTC CTT CCT CGA GTT GTC GGT CGT Ala Pro Val Gln Glu Gly Val Gln Gln Glu Gly Ala Gln Gln Pro Alas 690 700 CCT GCT ACA GCA CCT TCT CAA GGC GGA GTT AAT TCT CCT GTT AAT GTT GGA CGA TGT CGT GGA AGA GTT CCG CCT CAA TTA AGA GGA CAA TTA CAA Pro Ala Thr Ala Pro Ser Gln Gly Gly Val Asn Ser Pro Val Asn Val> 750 760 ACA ACT ACA GTT GAT GCT AAT ACA TCA CTT GCT AAA ATT GAA AAT GCT TGT TGA TGT CAA CTA CGA TTA TGT AGT GAA CGA TTT TAA CTT TTA CGA Thr Thr Thr Val Asp Ala Asn Thr Ser Leu Ala Lys Ile Glu Asn Alas 770 780 810 ATT AGA ATG ATA AGT GAT CAA AGG GCA AAT TTA GGT GCT TTC CAA AAT TAA TCT TAC TAT TCA CTA GTT TCC CGT TTA AAT CCA CGA AAG GTT TTA Ile Arg Met Ile Ser Asp Gln Arg Ala Asn Leu Gly Ala Phe Gln Asn>

TAA

101/133

850 860 830 840 820 AGA CTT GAA TCT ATA AAG AAT AGT ACT GAG TAT GCA ATT GAA AAT CTA TCT GAA CTT AGA TAT TTC TTA TCA TGA CTC ATA CGT TAA CTT TTA GAT Arg Leu Glu Ser Ile Lys Asn Ser Thr Glu Tyr Ala Ile Glu Asn Leu> 900 890 **-910**. 880 870 AAA GCA TCT TAT GCT CAA ATA AAA GAT GCT ACA ATG ACA GAT GAG GTT TTT CGT AGA ATA CGA GTT TAT TTT CTA CGA TGT TAC TGT CTA CTC CAA Lys Ala Ser Tyr Ala Gln Ile Lys Asp Ala Thr Met Thr Asp Glu Val> ~~~ ~950 930 940 " 920 GTA GCA GCA ACA ACT AAT ATG ATT TTA ACA CAA TCT GCA ATG GCA ATG CAT CGT CGT TGT TGA TTA TAC TAA AAT TGT GTT AGA CGT TAC CGT TAC Val Ala Ala Thr Thr Asn Met Ile Leu Thr Gln Ser Ala Met Ala Met> 990 970 980 ATT GCG CAG GCT AAT CAA GTT CCC CAA TAT GTT TTG TCA TTG CTT AGA TAA CGC GTC CGA TTA GTT CAA GGG GTT ATA CAA AAC AGT AAC GAA TCT Ile Ala Gln Ala Asn Gln Val Pro Gln Tyr Val Leu Ser Leu Leu Arg> 1010

FIGURE 40 (3 of 3)

#### Alignment List

Search Analysis for Sequence: B31-41kD Search from 1 to 1011 where origin = 1

Date: October 22,1993

Time: 15:03:24

Matrix: DNA database matrix Score Region from 1 to 1011

Maximum possible score: 4044

Database: Us		older	:: 41	kD	Flag	gell:	in c	lones	5							
+2 4 <b>.2</b> *		*	1	LO •	*		20		•	30	•	•	•	40	ŧ	
B31-41kD	ATG TAC	TTA AAT	ATC TAG	TAA ATT	CAT GTA	TTA ATT	ACA TGT	TCA AGT	GCT CGA	TAA	TAA -ATT	GCT -CGA-	TCA AGT	AGA TCT	AAT TTA	TAA ATT
1. KA-41kI [ 3996 ]				10			20			30				10		
2. P-Gau-4														-	•.••	••••
[ 3696 ]											• • •	• • •		10	•••	>
3. BO-41kI				LO			20			30				10		
[ 3684 ]	• • •	• • •	•••	• • •	• • •	• • •	•••	•••	• • •	• • •	• • •	•••	•••	• • •	• • •	>
4. DK29-41 [ 3672 ]			_	10			20			30				10		>
5. PKO-41)			1												•••	
[ 3672 ]	•••	•••									•••	•••		10	• • •	>
•	50		*	60		*	•	70	•		80		• ,	90		
B31-41kD	GGC CCG	TTA AAT	AAC TTG	GCT CGA	GCT CGA	AAT	CTT		AAA	ACT	CAA	GAA	AAG	CTT	TCT AGA	AGT TCA
1. KA-41ki [ 3996 ]	050			60				70			80			90		
												•••	•••		•••	
2. P-Gau-4	.c.		t			• • •		70		с	80	g		90		>
3. BO-41ki [ 3684 ]	D <b>50</b>			60				70			80			90		
[ 3684 ]	.c.	•••	t	• • •	•••	•••	•••	•••	•••	с	• • •	g	• • •	• • •	•••	>
4. DK29-4: [ 3672 ]			t	60				70	•		80	· a		90		>
5. PKO-41											•		•••		•••	•••
[ 3672 ]	.c.		t	•••		• • •	• • •	•••	• • •	с	80					>
•																
	1	00			110		.*	120		. *	1	30		3	L40 ★	
B31-41kD	CCC	TAC ATG	AGA TCT	TTA	AAT	CGA	GCT		GAT	GAT	GCT	GCT	GGC	ATG	GGA	GTT CAA

1. KA-41kD [ 3996 ] .		110		120	130		140
[ 3996 ] .	.g	•••	•••		130		140
2. P-Gau-4 [ 3696 ] .	.t	• • • • • •	• • • • • • •	••• ••• •	•• •••	t	g>
3. BO-41kD [ 3684 ]	100	110	•••	120	130	t	140 g>
4. DK29-41	100	110	a	120		t	g>
5. PKO-41k [ 3672 ]		110		120 '	130	t	140
	150	. 16	50	170	180	•	190
		a	COT CAA	אמג גדג (	GT TTG TCA CCA AAC AGI	CAA GC	TCT AGA A AGA TCT
1. KA-41kD [ 3996 ]		10	:	•••	· · · · · · · · · · · · · · · · · · ·		···>
2. P-Gau-4 [ 3696 ]	150 c.	1	60 	170 c.	180 ca	) 	>
3. BO-41kD	150 c.	1	60	170	180 ca		190 >
4. DK29-41	150 g .		,.	170	`a	o - · · · · ·	190 >
5. PKO-411 [ 3672 ]	k 150			170	18 ca	o 	190
	20	0			20		
B31-41kD			AA TTA T IT AAT A	TTA TTT T. AAT AAA A'	CAG ACA AC	A GAA GO T CTT CO	C TTA AAT
1. KA-41k [ 3996 ]	D 20	00	210		20 	230	240 >
2. P-Gau- [ 3696 ]	·4 20	00 a	210 c .		20	230 	240 .a >
3. BO-41) [ 3684 ]	сD 2	00 a	210 c .		20	230	.a>
4 27/20	41 2	00	210	2	220	230	240 .ag>
	1k 2	200	210	•	220	230	240 a>

				2:	50		;	260	•	_	270		_	2	80		
B	31-41kD	TAA ATT	GAA	GTA	GAA	AAA	GTC	TTA	GTA	AGA	ATG	AAG	GAA	TTG	GCA	ىلىك	CAA GTT
1	1. KA-41 3996 ]	kD		2	50			260			270			2	80		
	2. P-Gau 3696 ]	-4		2	50			260			270			2:	RO		
	3. BO-41 3684 J																
	4. DK29-																
	5. PKO-4 3672 ]	1k		25	50		:	260	•		270			21	RA		
		290		*	300			3:	10	•	3	320	•	* 1	330		
В	31-41kD	TCA AGT	GGT	AAC	GGC	ACA	TAT	TCA	GAT	GCA	GAC	AGA	GGT	TYT	ልጥል	CAL	ינה ע
ſ	1. KA-411 3996 ]	k290			300		• • •	3:	10	• • ;•		320	• • •		330	•••	>
Į	2. P-Gau 3696 ]	-290 	• • •		300 a	g		3:	10 c	• • •		320		•••	330	g	>
	3. BO-411 3684 ]				300 a	g		3:	10 c	t		320		•••	330	g	>
Į	4. DK29	4290			300 t	• • •		3:	10 c	•	• 3	320		• • •	330		>
Į	5. PKO-4: 3672 ]	1290	• • •		300 a	g	•••	3:	10 c	t		320	• • •		330	g	>
		34	10		3	350		•	360		•	37	0		3	80	
B.	31-41kD	GAA CTT	ATA TAT	GAG CTC	CAA GTT	CTT GAA	ACA TGT	GAC	GAA	TTA	TAA	AGA	TTA AAT	GCT CGA	GAT CTA	CAA GTT	GCT CGA
Į	1. KA-41 3996 ]	kD 3	10	• • •		350		• • •	360	• • •	•••	37	70			80	>
Į	2. P-Gau 3696 ]	-4 3	40	•••		350	• • •		360	• • •		37	70			g	>
ĺ	3. BO-41 3684 }	kD 3	40			350	• • •		360	• • •		37	70	• • •		g	>
	4. DK29-	41 3	40		:	350			360			. 37	70		3	80	

 $\{\hat{r}_i\}$ 

Į	3672 ]			•••		•••		•••	•••		•••	• • •	•••	• • •	• • •	•••	>
	5. PKO-411	c 34	0		3	50			360			37	0		3	80	
Į	3672 ]	•	•••			• • •				• • •	• • •	• • •	• • •	• • •	• • •	∙•g	>
			390			40	0		4	10		_	420			43	0
	31-41kD	* C>>	*	220	CNN	λTYC	C)C	* 27P4	TTA	TCA	AAC	AAA	TCT	GCT	TCT	CAA	TAA
В.	31-41KD	GTT	ATA	TTG	GTT	TAC	GTG	TAC	TAA	AGT	TTG	TTT	AGA	CGA	AGA	GTT '	TTA
	1. KA-41ki								•			•					
ſ	1. KA-41ki 3996 _F .)	D	390			41	•••			•••		:			• • •	•••	>
٠	2. P-Gau-									٠.	•		420			43	^
ſ	2. P-Gau-	4	390			4(	0		9	110		₹	**.		с		>
ı																	
	3. BO-41k	D	390			40	00		a	110			420		с	43	0 >
	4. DK29-4 3672 ]	1	390			4(	00		4	110			420		. C	43	0 >
	5. PKO-41	k	390			41	00	•	~	110			420		c	43	0
ι	5. PKO-41 3672 }	• • •	• • •	• • •	7	• • •	•••	• • •	9	•••	• • •	•••	•••	•••	•••	•••	• • • •
							450				<b>6</b> 0		,	30			480
																•	
B	31-41kD	GTA	AGA	ACA	GCT	GAA	GAG	CTT	GGA	DTA	CAG	CCT	GCA	AAA	TTA	AAC	ACA
		CAT	TCT													TTG	
	1. KA-41k	æ		440			450			4	60		4	170			480
I	3996 ]	• • •	• • •	• • •	• • •	••.•	• • •	• • •		• • •	•••	••:	•••	• • •	• • •	• • •	>
	2. P-Gau-	-4		440			450			4	60		4	170			480
1	3696 ]	• • •	.a.		• • •	• • •	• • •	• • •	• • •	• • •	•••	•••	• • •	• • •	• • •	• • •	>
	3. BO-41)	D D		440			450			4	60		•	470			480
1	[ 3684 ]	• • •	.a.	• • •	• • •		•••	• • •	• • •	• • •	• • •	• • •	• • •	• • •	• • •	•••	>
	4. DK29-4	41		440			450	)		4	60			470			480
	[ 3672 ]	• • •					a	٠	• • •	• • •	a	٠	• • •	• • •	с	•••	>
	5. PKO-4	1k		440			450	)		4	60			470			480
	[ 3672 ]	• • •	a.					•••		• • •	• • • •		• • •	• • •	•••	• • •	>
										•							
					490		ŧ	500			510	)	•	5	20	*	
	B31-41kD	CC	A GC	A TC	A CT	T TC	A GGC	CTI	CA)	A GC	G TC	r TGC	ACT	ATT	AGA	GTT	CAT
		GG	T CG	r ag	T GA	a ag	r ccc	GA2	A GT	r cc	C AG	A ACC	TGA	raa .	TCT	, CYY	GTA
	1. KA-41	kD			490			500			51	0		5	520		
	[ 3996 ]													• • • •			>
	2. P-Gau	-4			490			500			51	0		9	520		
	1 3696 1	•															>

				, , ,			
3. BO-41ki [ 3684 ]	·	490	a	500 tc	510	•••	520
4. DK29-41 [ 3672 ]	ι g	490	a	500 tc	510 t		520
5. PKO-411 [ 3672 ]		490	a	500 tc	510 t	••• •••	520 >
· .	5 <b>30</b>	540	•	550	5	60 . *	570
					TAA CGA		ATT TAT GCA GCT TAA ATA CGT CGA
1. KA-41k [ 3996 ]	530	540 a.	•••	550 		60 ••• •••	570 >
2. P-Gau-	530 g	540 at		550 a		60	570 t>
3. BO-41ki						60	570 t>
4. DK29-4 [ 3672 ]	530 g	540 at		550 g		60	570t>
5. PKO-41 [ 3672 ]							570 t>
	580	<u>.</u>	590 *	600	•	610	620
B31-41kD		GCA AAT	CTT TTC	TCT GGT	GAG GGA	GCT CAA	ACT GCT CAG GCT IGA CGA GTC CGA
1. KA-41k [ 3996 ]							620
							620 g >
							620 g >
4. DK29-4 [ 3672 ]	1 580		590 a	600		610 g	620 g a>
5. PKO-41 [ 3672 ]				600 g.,		610	620 g >
	630	)	640	•	650	660	670
B31-41kD							CAA CAG CCA GCA GTT GTC GGT CGT

									·	_							
ι	1. KA-41ki 3996 }	· · · ·	630		•••	64	0	•••		550			660	•••	•••	67	70 >
Į	2. P-Gau-4	• • •	630 t			64	0	.c.		550 g.a	•••	•••	660	g	a	67	70 a>
Į	3. BO-41kI		630 t			64	0	.c.		550 g.a	•••	•••	660	g	a	6	70 a>
	4. DK29-41																
·[	5. PKO-41)	· · · ·	630 t	•••	•••	64	0	.c.		550 g.a	` .	•••	660	g	a	67	70 a>
				680		•	690			70				710	•	٠.	720
ВЗ	31-41kD				GCA		TCT	CAA	GGC	GGA	GTT	TAA	TCT	CCT	GTT		GTT
	1. KA-41ki 3996 ]																
	2. P-Gau-																
	3. BO-41ki 3684 ]																720 >
	4. DK29-4: 3672 ]																
	5. PKO-41 3672 ]																
				7	30	*		740			750			76	60		
B	31-41kD			ACA	GTT		GCT	AAT	ACA	TCA	CTT	GCT	AAA	ATT	GAA	AAT	
	1. KA-41k 3996 ]	D 			30	•••		740	•••	•••	750 •••	•••	•••	7	60		>
Į	2. P-Gau- 3696 ]				30			740					•••		60		>
Į	3. BO-41k	 Φ	c	7	30	•••	•••			•		• • •	• • •		60		>
I	4. DK29-4 3672 ]									t	750				60		>
[	5. PKO-41	.k 	(	7	30	• • •	• • •	740		• • •	750 	•••	•••	7 a	60		>

								7.								
	770			780		•	79	90	•	8	800 *		•	810		•
B31-41kD	ATT AAT	AGA	ATG	ATA	AGT	GAT	CAA	AGG	GCA	TAA	TTA	GGT	GCT CGA	TTC AAG	CAA GTT	TAA ATT
1. KA-41) [ 3996 ]	c770	•••	•••	780	•••	•••	79	90	• • •		300	•••	•••	810	•••	>
2. P-Gau- [ 3696-]	. •••	• • •	• • •	• • •	• • •	• • •	• • •	a	•••	•••	•••	•••	•••	810	•••	>
3. BO-411 [ 3684 ]	• • •	• • •	• • •	•••	• • •	• • •	•••	a	•••	•••	•••	• • •	•••		•••	>
4. DK29-	• • •	•••	•••	•••	•••	• • •	•••	a	•••	• • •	•••	•••	•••	810	·••	>
5. PKO-4	1770	•••	•••	780 ::·	• • •	•••	7:	90 a	•••	•••	300	•••	•••	810	•••	>
	82	20		1	830		•	840		*	85	50		8	60	
B31-41kD	AGA TCT	CTT GAA	GAA	TCT	ATA	AAG	TAA	AGT	ACT	GAG	TAT	GCA	ATT		TAA	CTA GAT
1. KA-41 [ 3996 ]	kD 82	20			830	•••		840	• • •	•••	8:	50	•••		60	>
2. P-Gau [ 3696 ]	-4 83	20	•••	• • •	830	• • •		840 c		•••		50 t				>
3. BO-41 [ 3684 ]	• • •	•••	• • •	•••	• • •	•••	• • •		• • • •	• • •	•••	t	•••		• • •	>
4. DK29- [ 3672 ]	• • •	• • •	9		• • •	• • •	g	• • •	• • •	•••	• • •	t	• • •	• • •	с	>
5. PKO-4	11k 8	20	• • •	•••	830	• • •	• • • •	840	) :	•••		50 t	•••		360	>
	•			•	8	-		t						4		•
B31-41kD														CTA		
1. KA-41 [ 3996 ]	lkD 								890		•	900		•••	-	10
2. P-Gar [ 3696 ]									890					•••		10
3. BO-4 [ 3684 ]									890						-	10
4. DK29 [ 3672 ]									890						-	10>

5. PKO-41)	k	870			88	30		ε	90			900				
[ 3672 ]	• • •	•••	•••	•••	• • •	• • •		•••	•••	•••	•••	•••	•••	•••	•••	•••>
	*	9	920		•	930		*	94	10	*	9	950		•	960
B31-41kD	GTA CAT	GCA CGT	GCA CGT	ACA TGT	ACT TGA	TAA ATT	ATG TAC	TTA AAT	ATT TAA	ACA TGT	CAA GTT	TCT AGA	GCA CGT	ATG TAC	GCA CGT	ATG TAC
1. KA-41kl	D 		920	•••		930								•••		960 •••>
2. P-Gau-	4		920 t	•••	•••	930	.gt	•••	94	10` . <u>.</u> t	•	::	950	•••	•••	960 >
3. BO-41ki [ 3684 ]	D 		920 t		•••	930	.gt		94	40 t			950	•	·.	960 · >
4. DK29-4 [ 3672 ]	1		920 t	• • •		930 	.gt		9	40		ġ.	950		• • •	960 >
5. PKO-41 [ 3672 ]	k 	•••	920 t	•••		930 a	.gt	•••	9.	40 t	•••	•••	950	•••		960 >
		•	9	70	*		980		•	990		*	10	00	*	
B31-41kD	TTA AAT	. ccc	CAG	GCT CGA	TAA	CAA	GTT	CCC	CAA	TAT	GTT	TTG AAC	TCA AGT	TTG AAC	CTT GAA	AGA TCT
1. KA-41k [ 3996 ]	 D		9	70			980			990		•••				>
2. P-Gau- [ 3696 ]			و	70			980	t		990		• • •	10	00		>
3. BO-417 [ 3684 ]	മ		2	970		. <b></b> .	980	t				•••		00		>
4. DK29-4 [ 3672 ]	11			970 a			980	t		990	•		10	00	•••	
5. PKO-41	lk 		a	970			980	t	·	990		• • •	10	00		>
:	1010								•							
B31-41kD	AT TA												٠.			
2. P-Gau [ 3696 ]		.>														

Sequence Rar	nge: 1 t	o 822		•				
		10	•	20	30	•	40	•
OspA-B31	ATG AAA TAC TTT	አአአ ጥልጥ	TTA TTG	GGA ATA	GGT CTA	ATA TTA TAT AAT	GCC TTA CGG-AAT	ATA GCA TAT CGT
OspA-B31 [ 3288 ]		10		20	30	•	40	>
OspA-KA [ 3288 ]		10	•••	20	`30	•••	40	••• •••>
OspA-N40 [ 3276 ]	•••			20	30	•••	40	··>
OspA-ZS7 [ 3264 ]				20	30	• • • • • •	40	>
OspA-25015 [ 2802 ]				20	30	•••	t	>
OspA-TRO [ 2648 ]		10		20	30	•••	40	>
OspA-K48 [ 2584 ]		10		20	30  30	•••	40	>
OspA-HE 11 [ 2580 ]	•••	10		20  20	30	•••	40	>
OspA-DK29 [ 2566 ]	•••	10	••••	20		• • • • • •	40	>
OspA-Ip90 [ 2562 ]	•••			20		•••	a	>
OspA-BO [ 2558 ] OSPA-IP3		10 	•••	20		•••	40	>
[ 2558 ] OspA-PKO	•••		•••	20		•••	40	>
[ 2558 ] Ospa-acai	•••	10	• • • • • • •	20		 )	40	>
[ 2556 ] ospa-P-GAU	•••	10		20		)	40	>
[ 2544 ]	•••	• • • • •		• ••• ••		• • • • • •		>
	50 •	*	• •	70	•	80	90	•
OspA-B31	TGT AZ ACA TI	G CAA AA' C GTT TT	T GTT AG A CAA TC	C AGC CT G TCG GA	T GAC GAG A CTG CTG	)AA AAA C YIT TIT C	AGC GTT TCG CAA	TCA GTA AGT CAT

FIGURE 42 (1 of 16)

: 4

111/133

OspA-B31 [ 3288 ]	50	60	70	80	90
OspA-KA [ 3288 ]	50	60	70	80	90
OspA-N40 [ 3276 ]	50	60	70	80	90>
OspA-ZS7 [ 3264 ]	50	60	70 	80	90>
OspA-25015 [ 2802 ]	50	60	70 '	80	90
OspA-TRO [ 2648 ]	50	60	70 t	80	.90
OspA-K48 [ 2584 ]	50	60	70 ta	80 t	90
OspA-HE 11 [ 2580 ]	50	60 	70 ta	80t	90>
OspA-DK29 [ 2566 ]	50	60	70 ta	80 t	90>
OspA-Ip90 [ 2562 ]	50	60	70 ta	80 t	90>
OspA-BO [ 2558 ]	50	60	70 ta	80	90 .c>
OSPA-IP3 [ 2558 ]	50 c	60	70 ta		90 .c>
OspA-PKO [ 2558 ]	50 c	60	70 ta		90 .c>
OspA-ACAI [ 2556 ]	50 c	60	70 ta	80	90 .c>
ospA-P-GAU [ 2544 ]	50 c	60	70 te	80	90 .c>
	100	110	120	130	140
OspA-B31	GAT TTG CCT CTA AAC GGA	CCA CTT TAC	AAA GIT CIT GIA TII CAA GAA CA	AAD AAA CDA A TTC TTT DOT 1	TTT TTT TTT
OspA-B31 [ 3288 ]		110			140>
OspA-KA [ 3288 ]	100	110	120	130	
Ospa-N40	100	110	120	130	140

FIGURE 42 (2 of 16)

SUBSTITUTE SHEET (RULE 26)

		•			
[ 3276 ]			.c		>
OspA-ZS7 [ 3264 ]	100		120 .c	130	140>
OspA-25015 [ 2802 ]	100	110	120	130	140 g>
OspA-TRO [ 2648 ]	100 a		120	130	140 g>
OspA-K48 [ 2584 ]	100 a	110g	120 c	· 130	140 g>
OspA-HE 11 [ 2580 ]	100a	110 g	120	<del>-1</del> 30 t	140 g>
OspA-DK29 [ 2566 ]	100 a	110 g	120 c	130 t	140 g>
OspA-Ip90 [ 2562 ]	100 a	110 g c	120	130°	140 g>
OspA-B0 [ 2558 ]	100	110 g	120	130 t	140 g>
OSPA-IP3 [ 2558 ]	100	110 gt .	120		140 g>
OspA-PKO [ 2558 ]	100	110 g	120	130 t	140 g>
Ospa-ACAI [ 2556 ]	100	110 g	120 '	130 t	140 g>
ospA-P-GAU [ 2544 ]	. 100	110 g	120	130 t	140 g>
	150	160	170	180	190
OspA-B31	GAC GGC AAG CTG CCG TTC	TAC GAT CTA A ATG CTA GAT T	TT GCA ACA GTA AA CGT TGT CAT	GAC AAG CTT CTG TTC GAA	GAG CTT AAA CTC GAA TTT
OspA-B31 [ 3288 ]	150	160	170	180	190
OspA-KA [ 3288 ]	150	160		180	190
OspA-N40 [ 3276 ]	150	160	170	180	190
OspA-ZS7 [ 3264 ]	150	160	170	180	190 >
OspA-25015 [ 2802 ]	150	160 ag	170	180	190

FIGURE 42 (3 of 16)

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113/133

•															•	
OspA-TRO [ 2648 ]	t	150 t	a	•••	16 ag.	0 • • • •	g		70	•••	•••	180	a	•••		90
OspA-K48 [ 2584 ]	•••	150 t	a	• • •	16 ag.	0 • • • •	gag		70	•••	•••	180		•••		•••>
OspA-HE 11 [ 2580 ]	t	150 t	a	•••	16 ag.	0	g			•••	a	180	•••	•••		90 •••>
OspA-DK29 · [ 2566 ₇ ]	•••	150 t	a	•••	16 ag.	0	gag		.70	•••	:		•••	•••		30 •••>
OspA-Ip90 [ 2562 ]	t	150 t	a		16 ag.	0	g		.70	•••	•••	180	•••	•••		90 •••>.
OspA-BO [ 2558 ]	•••	150 t		•••	16 ag.	0	.ag		.70	•••	•••				19 a	0 •••>
OSPA-IP3 [ 2558 ]	· • •	150 t			16 ag.	0	.ag		.70	•••	•••			•••	19 a	>
OspA-PKO · · · [ 2558 ]	•••	150 t	•••		16 ag.		.ag		.70				a	•••	19 a	
Ospa-ACAI [ 2556 ]	•••	150 t	•••		16 ag.		.ag	•:•		•••	•••	180	a	•••	19 a	
ospA-P-GAU [ 2544 ]	•••	150 t	•••	•••	16 ag.		.ag			•••	•••	180		•••		>
		2	200			210				20		:	230		_	240
OspA-B31	GGA CCT	ACT TGA	TCT AGA	GAT CTA	AAA TTT	244	AAT	GGA	TCT	GGA	GTA CAT	CTT GAA	GAA CTT	CCC	GTA CAT	AAA TTT
OspA-B31 [ 3288 ]	•••		200			210	•••			20	•••		230		•••	240 >
Ospa-Ka [ 3288 ]			200			210				20		•••	230		•••	240
OspA-N40 [ 3276 ]	•••		200			210	•••			20	•••		230	•••	•••	240 >
OspA-ZS7 [ 3264 ]	•••	• • •	200			210				20	• • •	•••	230	•••	•••	240
OspA-25015 [ 2802 ]	•••		200			210				20 g	••9	• • •	230		•••	240
Ospa-TRO [ 2648 ]		•••	200			210 .g.		:t		220	ac.	• • •	230	t	.a.	240 >
OspA-K48 [ 2584 ]			200			210		:t		220	ac.		230		.a.	240

FIGURE 42 (4 of 16)

										20		240
OSPA-HE 11 [ 2580 ]		200		210	c .	.t		ac			.t .a.	240
OspA-DK29 [ 2566 ]		200		210	c .	.t	220	ac		230	.t .a.	240 >
OspA-Ip90 [ 2562 ]		200		210	c .	.t	220	ac	:	230	.t .a.	240 •••>
OspA-BO [ 2558 ]		200		210 g	••••	.t	220	.g	g	230	.t ac.	240 >
OSPA-IP3 [ 2558 ]		200		210 g	•••	.t	220			230	.t ac.	240 >
OspA-PKO [ 2558 ]		200		210 g		.t	220			230	.t ac.	240
OspA-ACAI [ 2556 ]	•••	200		210 g		t	220				.t ac.	240 >
ospA-P-GAU		200		210 g		t	220			230	.t ac.	240 >
, 2517 ,		25			260		7	270		280		
OspA-B31	GCT C	25 GAC AAA CTG TTT	AGT AAA	ста	AAA 1	ATA TAA	ACA A	· ATT TO	T GAC	GAT CTA G	TA GGT	CAA GTT
OspA-B31 [ 3288 ]	• • •	25	0		260			270		280		>
OspA-KA [ 3288 ]	•••				260	• • •		270 · · · · ·	• • • • • • • • • • • • • • • • • • • •	280		>
	•••	25 25		•••		•••		270  270 		280  280	•••••	>
[ 3288 ] Ospa-N40	•••	25 		•••	 260			270 270 270	•••••	280		>
[ 3288 ] OspA-N40 [ 3276 ] OspA-ZS7	•••	25	50	•••	260 260 260	•••		270 270 270 270	••••	280  280 		>> ac.>
[ 3288 ] Ospa-N40 [ 3276 ] Ospa-ZS7 [ 3264 ] Ospa-25015	  	25	50  50 	• • • • • • • • • • • • • • • • • • • •	260 260 260 260	•••		270 270 270 270 270 g	•••••	280  280 	  ) a.c	
[ 3288 ]  Ospa-N40 [ 3276 ]  Ospa-ZS7 [ 3264 ]  Ospa-25015 [ 2802 ]  Ospa-TRO	  t	25	50  50 	• • • • • • • • • • • • • • • • • • • •	260  260  260  260	•••		270 270 270 270 270 g		280 280 280 280		a>
[ 3288 ] OSPA-N40 [ 3276 ] OSPA-ZS7 [ 3264 ] OSPA-25015 [ 2802 ] OSPA-TRO [ 2648 ] OSPA-K48	a	29	50 50 50 50 50		260 260 260 260 260	•••		270 270 270 270 270 270 270 270	t	280 280 280 280 280		a>
[ 3288 ] OSPA-N40 [ 3276 ] OSPA-ZS7 [ 3264 ] OSPA-25015 [ 2802 ] OSPA-TRO [ 2648 ] OSPA-K48 [ 2584 ] OSPA-HE 11	a	25 	50 	· · · · · · · · · · · · · · · · · · ·	260  260  260  260 	•••		270 270 270 270 270 270 270 270 270 270		280 280 280 280 280 280	a.c	a> >

FIGURE 42 (5 of 16)

				•												
[ 2562 ]	a	•••	• • •	• • •	• • •	.c.	• • •			•••	g.,	9	• • • •		a.,	a>
OspA-BO [ 2558 ]	.a.	•••		50	•••		260 	•••	•••	270	g	•••		80	•••	a>
OSPA-IP3 [ 2558 ]	.a.	•••	2	50	•••		260 	•••	•••	270				80	a	a>
OspA-PKO [ 2558 ]	.a.	•••		50	•••		260		•••	270		•••		80	•••	a>
OspA-ACAI [ 2556 ]	.a.			50			260		•••			•••		BO	a.,	a'>
ospa-P-GAU [ 2544 ]	.a.	•••	29	50			260			270		•••	21		a	a>
	290		_	300		•	3:	10	•		320			330		
OspA-B31	ACC TGG	ACA TGT	CTT GAA	GAA	GTT	TTC	AAA	GAA	GAT	GGC	AAA	ACA	CTA	GTA	TCA AGT	AAA TTT
OspA-B31 [ 3288 ]	290	•••	• • •	300					•••		320			330	• • •	>
OspA-KA [ 3288 ]	290	•••												330		>
OspA-N40 [ 3276 ]	290	•••		300										330		>
OspA-ZS7 [ 3264 ]	290		• • •								320		•	330	•••	>
OspA-25015 [ 2802 ]			• • •	300		a	31				320		t	330 g	• • •	>
OspA-TRO [ 2648 ]	290			300			31	10		:	320			330		
Ospa-K48 [ 2584 ]	290 t		t		a						320	•••	t	330	• • •	>
OspA-HE 11 [ 2580 ]	290			300							320		t	330	g	>
OspA-DK29 [ 2566 ]				300			31	LO			320			330		>
OspA-Ip90 [ 2562 ]	290			300			31	LO		;	320			330		>
OspA-BO [ 2558 ]	290			300			31	10		:	320			330		.g.>
OSPA-IP3 [ 2558 ]	290			300			3:	10		:	320			330		.g.>
	•					- • •	- • •				• • •	• • •		• • 9	• • •	.9

FIGURE 42 (6 of 16)

OspA-PKO		300	310	320	330
[ 2558 ]	t.c				tgg.>
OspA-ACAI [ 2556 ]	290 t.c	300 c	310 	320	330 tgg.>
ospA-P-GAU [ 2544 ]	290 t.c	300 c	310 a	320	330 tgg.>
	340	350	360	370	380
OspA-B31	AAA GTA ACT TTT CAT TGA	TCC AAA GAG AGG TTT CTG	C AAG TCA T G TTC AGT A	CA ACA GAA GAA GT TGT CTT-CTT	AAA TTC AAT GAA TTT AAG TTA CTT
OspA-B31 [ 3288 ]	340	350	360 	370	380
OspA-KA [ 3288 ]	340	350	360	370 .	380
OspA-N40 [ 3276 ]	340	350		370	380
OspA-ZS7 [ 3264 ]	340	350	360		380
OspA-25015 [ 2802 ]	340 ag	350	360 t		380
OspA-TRO [ 2648 ]	340 a.	350 t		370t	380 c .c.>
OspA-K48 [ 2584 ]	340	350 ctt	360	370	380
OspA-HE 11 [ 2580 ]	340	350 ctt	360	370	380 c>
OspA-DK29 [ 2566 ]	340	350		370	380c .g.>
OspA-Ip90 [ 2562 ]				370	380 c .c.>
OspA-BO [ 2558 ]	340g	350 t	360 a a	370 t	380
OSPA-IP3 [ 2558 ]	340 g	350 t	360 a a	370 t	.tg>
OspA-PKO . [ 2558 ]		350 t	360	370 t	.tg>
OspA-ACAI [ 2556 ]	340 g	350 t	360	370 t	380 .

FIGURE 42 (7 of 16)

ospA-P-GAU [ 2544 ]	340 35	0 ga	360 a	370	380 J>
OspA-B31	390 AAA GGT GAA GTA T TTT CCA CTT CAT A	KAA AAN 100	አጥኔ ኔጥኔ ልርል	AGA GCA GA	C GGA ACC AGA
OspA-B31 [ 3288 ]	390	400	410	420	430
OspA-KA [ 3288 ]			•••	•••	• • • • • • • • • • • • • • • • • • • •
OspA-N40 [ 3276 ]	390	•• •••	•••	•••	
OspA-ZS7 [ 3264 ]	390	••••••	410	••• •••	430 
OspA-25015 [ 2802 ]	390 c t 9	jt	g g	a.	430
Ospa-TRO [ 2648 ]	t	400	.c ct.	a.	g>
Ospa-K48 ( 2584 ) Ospa-HE 11	g ac.	400	.c gt.	420	430
[ 2580 ] Ospa-DK29	390 g a 390 g ac.	400	410	420	430
[ 2566 ] OspA-Ip90 [ 2562 ]		400	410	420	430
OspA-BO [ 2558 ]	390	400	410	420	430
OSPA-IP3 [ 2558 ]	t.g	c	.ccg		
OspA-PKO [ 2558 ]	t.g	c	ccg	420 a. a 420	
OspA-ACAI [ 2556 ]	t.g		ccg	a. a 420	.ta.>
ospA-P-GAU [ 2544 ]	t.g	c	ccg	a. a	.t ,a.>
	440	450 • •	460		• •
OspA-B31	CTT GAA TAC ACA	GGA ATT AA	A AGC GAT GG	ia toți gga a	AA GCT AAA GAG

FIGURE 42 (8 of 16)

Assaurter Aller Inile (A)

	GAA CTT ATG TGT	CCT TAA TTT	TCG CTA CCT	AGA CCT TTT	CGA TTT CTC
OspA-B31 [ 3288 ]	440	-	460	470	480
OspA-KA [ 3288 ]	440		460	470	480
OspA-N40 [ 3276 ]	440		460	470	480
OspA-ZS7 [ 3264 ]	440		460	470	480
OspA-25015 [ 2802 ]	440		460	470	480 a>
OspA-TRO [ 2648 ]	440	450 .aa	460	470 a.c	480 a>
OspA-K48 [ 2584 ]	440		460		480 a>
OspA-HE 11 [ 2580 ]	440	450 .aca	460 aa.	470 a.c	480 a>
OspA-DK29 [ 2566 ]	440	450 .aca	460	470 c	480 a>
OspA-Ip90 [ 2562 ]	440		460 aa.	470 a.c	480 a>
OspA-BO [ 2558 ]	440 t		460		480 د
OSPA-IP3 [ 2558 ]	440 t	450 .ag	460	470 a.c	480 د
OspA-PKO [ 2558 ]	440 t		460		480 a>
OspA-ACAI [ 2556 ]	440 t			470 a.c	480 a>
ospA-P-GAU { 2544 }	440t			470 a.c	
	490	500	. 510	520	•
OspA-B31	GTT TTA AAA GGC CAA AAT TTT CCG	TAT GTT CTT ATA CAA GAA	GAA GGA ACT CTT CCT TGA	CTA ACT GCT G	AA AAA ACA TT TTT TGT
OspA-B31 [ 3288 ]	490	500	510	520	
Ospa-Ka { 3288 }	490	500	510	520	

Ospa-N40		490	500	510	520
[ 3276 ]			•••	••• ••• •••	>
OspA-ZS7 [ 3264 ]		490 . a	500	510 t	520
OspA-25015 [ 2802 ]	ac	490 aa	500	510	520 · · · · · · · g>
OspA-TRO [ 2648 ]	.c	490 at.	500 .c	510 g	520 cc>
					cgġ I
OspA-K48 [ 2584 ]		490 at.	500 ac	510 _. g	5201 530
					cgg I
OspA-HE 11 [ 2580 ]	••• ••••	490 at.	500 ac	510 ·	
					cgg I
OspA-DK29 [ 2566 ]		490 at.	500 ac	510 g	5201 530 >
				•	cgg 
OspA-Ip90 [ 2562 ]		490 at.	500 .c	510 g	5201 530>
OspA-BO [ 2558 ]		490 aat.	500 ac	510 aa g g	520 aat gt.>
OSPA-IP3 [ 2558 ]		490 aat.	500 ac	510 aa g g	520 aat gt.>
OspA-PKO [ 2558 ]		490 aat.	500 ac	510 aa g g	520 aat gt.>
Ospa-ACAI [ 2556 ]		490 aat.	500 ac	510 aa g g	520 aat gt.>
ospA-P-GAU [ 2544 ]		490 aag .t.		510 aa g g	520 aat gt.>
	530	540	550	. · · 560	570
OspA-B31	• •	•	• •	• •	AAA AAT ATT TCA
בנט ייגנט	TOT AAC C	AC CAA TTT	CTT CCT TGA	CAA TGA AAT TCC	TTT TTA TAA AGT
OspA-B31 [ 3288 ]	530	540	550 	560	570 >
Ospa-Ka	530	540	550	560	570

FIGURE 42 (10 of 16)

[ 3288 ]	•••		•••	•••		•••		• • •	• • • •	• • • •	• • • •		• • • •	• • • •	>
OspA-N40 [ 3276 ]	530		540							560		• • • •	570		>
OspA-ZS7 [ 3264 ]	530		540				50			560			570		>
OspA-25015 [ 2802 ]			540			5	50			560			570		
OspA-TRO	530		540			5	50		•	560			570		
OspA-K48 [ 2584 ]	•••	540 aaa		.c.						•••			c	٠.	.t.>
OspA-HE 11 [ 2580 ]	•••	540 aaa	•••	.c.	5: g	50 c			560	• • •	···	570	с	• • •	>
OspA-DK29 [ 2566 ]	•••	540			5:	50			560			570			
OspA-Ip90 [ 2562 ]	a	540 aaa				50 c			560 gt.		•••	570	c.c	•••	>
OspA-BO [ 2558 ]	530									560					g>
OSPA-IP3 [ 2558 ]	530	.aa	540 a			55				560		g	570 g.a		g>
OspA-PKO [ 2558 ]	530	.aa					50 c				t		570 g.a	•••	g>
OspA-ACAI [ 2556 ]	530	.aa	540 a		•••		50 c			560	t	g	570 g.a	•••	g>
ospA-P-GAU [ 2544 ]			540 a									g	570 g.a	•••	g>
	580		<b>Š</b> !	90			600		_	63	10		e	520	
OspA-B31	AAA TCI TTT AGA					GTT	GAA	CTT	AAT	GAC	ACT				
OspA-B31 [ 3288 ]	580	•••	5!	90			600	•••	-:-	61	10			20	>
OspA-KA [ 3288 ]	580	•••	55	90	•••	• • •	600	•••	•••	63		•••		20	>
OspA-N40 [ 3276 ]	580	•••	55	90		• • •	600			6:	10	•		520	>
OspA-ZS7 [ 3264 ]	580		5:	90			600			6	10		6	520	>

FIGURE 42 (11 of 16)

OspA-25015 [ 2802 ]	580	590 aa a	600 .c	610	620c. caa>
OspA-TRO [ 2648 ]	c	a a.a a	g	t	620 a tcc. cag>
OspA-K48 58	30	590 a a.a a	600 c	610 gt	620c. cag>
OspA-HE 1158	30 ċ	590 · a a.a a	600 c	610 g	620 tcg>
	c		•	gt	620 cccgg>
OspA-Ip90 58 [ 2562 ]	30 c	590 a a.a a	600 g	610 t	620 c. cag>
OspA-BO [ 2558 ]	580	590 aa a	600 ct	610	620 acc. cag>
OSPA-IP3 [ 2558 ]	580	590 aa a	600 ct	610	620 acc. cag>
OspA-PKO [ 2558 ]	580	590 aa a		610	620 acc. cag>
OspA-ACAI [ 2556 ]	580	590 aa a	600 ct	610	620 acc. cag>
ospA-P-GAU [ 2544 ]	580	590 aa a	600 ct	610	620 acc.cag>
·	630	640	650	660	670
OspA-B31	GCT ACT CGA TGA	AAA AAA ACT GCA TTT TTT TGA CGT	GCT TGG AAT CGA ACC TTA	TCA GGC ACT AGT CCG TGA	TCA ACT TTA ACA AGT TGA AAT TGT
OspA-B31 [ 3288 ]	630		650		670
OspA-KA [ 3288 ]	630 	640		660	670
OspA-N40 [ 3276 ]	630 				670
OspA-ZS7 [ 3264 ]	630	640	650	660	670
OspA-25015 [ 2802 ]	630		650 aaa g	660 g	670
OspA-TRO [ 2648 ]	630		650 aaa g	660 aat	670

FIGURE 42 (12 of 16)

OspA-K48 [ 2584 ]	630	640 	650 aaa g	660 aaa	670
OspA-HE 11 [ 2580 ]			650 a.a g	t	670
OspA-DK29 [ 2566 ]	630	640 • • • • • • • • • • • • • • • • • • •			670 .c>
OspA-1p90 [ 2562 ]	630	640 • • • • • • • • • • • • • • • • • • •	650 a.a g	660 aag	670 .c
OspA-BO [ 2558 ]	630	640	650	660	670
OSPA-IP3 [ 2558 ]	630	640	650	660	670 ····>
OspA-PKO [ 2558 ]	630	640	650	660	670 t>
Ospa-ACAI [ 2556 ]	630	640	650	660	670 t>
ospa-P-GAU [ 2544 ]	630	640	650	660	670 t>
OspA-B31	680 ATT ACT GTA TAA TGA CAT	690 • • • • • • • • • • • • • • • • • • •	700 AAA ACT AAA GA TTT TGA TTT CT	יייה יאנט זהנט ט יייי	T 202 222 022
OspA-B31 [ 3288 ]	680	690	700	710	
Ospa-Ka [ 3288 ]	680	690	700		
OspA-N40 [ 3276 ]	680	690	700	710	720
OspA-ZS7 [ 3264 ]	680	690	700	710	720
OspA-25015 [ 2802 ]	680	690 ac	700 c.	710 a	720 c>
OspA-TRO [ 2648 ]	680 gg	690	a	710 aa	720 
OspA-K48 [ 2584 ]	680	690	700 c a	710	720
OspA-HE 11 [ 2580 ]	680	690	700 c.a	710	720
OspA-DK29	680	690	700	710	720

FIGURE 42 (13 of 16)

[ 2566 ]	ggt			.ac >
OspA-Ip90 [ 2562 ]	680ggt	590 70 c cgc	00 71	.ac>
OspA-BO [ 2558 ]	680	690	700	
OSPA-IP3	680	690	700	710 720
[ 2558 ]	gt	•••••		t c>
OspA-PKO [ 2558 ]	680 gt	690 c	700 .c. c.a	710 720 c>
Ospa-ACAI [ 2556 ]	680 gt		700 .c. c.a	710 720
ospA-P-GAU [ 2544 ]	680 gt	690 c	700 .c. c.a	710 720 c>
	730	740	750	760
OspA-B31	AAC ACA ATT ACA O	TA CAA CAA TAC	GAC TCA AAT C	SEC ACC AAA TTA GAG
OspA-B31 [ 3288 ]	730		750	760
Ospa-Ka [ 3288 ]	730	740		760 >
OspA-N40 [ 3276 ]	730	740	750 、	760
OspA-ZS7 [ 3264 ]	730	740	750	760 >
OspA-25015 [ 2802 ]	730 g		750 gca	760 acga>
Ospa-TRO [ 2648 ]	730 ga	740 a	750 gca	760 t ca>
OspA-K48 [ 2584 ]	730 ga	740 a	750 gca	760 770 t ca>
OspA-HE 11 [ 2580 ]	730 ga	740 a.c	750 gca	760 t ca>
OspA-DK29 [ 2566 ]	730 ga	740 ag	750 gca	760 770t ca>
OspA-Ip90 [ 2562 ]	730 ga	.740 a	750 gca	760 · 770t ca>
OspA-BO [ 2558 ]	730 gat	740 a	750 c gca	760 t'ta>

FIGURE 42 (14 of 16)

OSPA-IP3	730 740 750 gatac gcat	760
[ 2558 .]		
OspA-PKO [ 2558 ]		ta>
OspA-ACAI [ 2556 ]		760 ta>
ospA-P-GAU [ 2544 _c ]		760 t _.
•	770 780 790 800	810
OspA-B31	GGG TCA GCA GTT GAA ATT ACA AAA CTT GAT GAA ATT AA CCC AGT CGT CAA CTT TAA TGT TTT GAA CTA CTT TAA TT	A AAC GCT TTA
OspA-B31 [ 3288 ]	770 780 790 800	810
OspA-KA [ 3288 ]	770 780 .790 800	, 810 
Ospa-N40 [ 3276 ]	770 780 790 800 	810
OspA-ZS7 [ 3264 ]		>
OspA-25015 [ 2802 ]	770 780 790 800c a ac	810
	c aaccac C	>
	c aacc a.a c	
OspA-HE 11 [ 2580 ]	770 780 790 800c aa c c a.a c	810
OspA-DK29 [ 2566 ]	c aacc a.a c	0>
OspA-Ip90 [ 2562 ]	780 790 800 81 c aaccg a.a c	0 . g.t>
OspA-BO [ 2558 ]	770 780 790 800c a c ac c	810>
OSPA-IP3 [ 2558 ]	770 780 790 800 c acac C	
OspA-PKO [ 2558 ]	770 780 790 800c a c	810>

FIGURE 42 (15 of 16)

Ospa-ACAI 770 { 2556 }c a	780 c	790 .ac	800 c	810 • • • • • • • • • • • • • • • • • • •
ospA-P-GAU 770 [ 2544 ]c a	780 c	790 .ac	800 c	810

820 AAA TAA OspA-B31 TTA TTT OspA-B31 820 [ 3288 ] ... ...> 820 Ospa-KA [ 3288 ] ... ...> Ospa-N40 820 [ 3276 ] ... ...> OspA-ZS7 820 [ 3264 ] ... ...> OspA-25015 [ 2802 ] .g.> OspA-TRO 820 [ 2648 ] ... ..> OspA-K48 820 [ 2584 ] ... ...> OspA-HE 11 820 [ 2580 ] ... ..> OspA-DK29 820 [ 2566 ] ... ...> OspA-Ip90 820 [ 2562 ] 820 OspA-BO [ 2558 ] ... ..> OSPA-IP3 820 [ 2558 ] ... ..> OspA-PKO 820 [ 2558 ] ... ..> OspA-ACAI 820 [ 2556 ] ... ...> ospA-P-GAU 820 [ 2544 ]

... ...>

30 ATG AAA AAA TAT TTA TTG OGA ATA GGT CTA ATA TTA GCC TTA ATA TAC TIT TIT ATA AAT AAC CCT TAT CCA GAT TAT AAT CGG AAT TAT 60 70 GCA TGT AAG CAA AAT GTT AGC AGC CTT GAT GAA AAA AAT AGC GTT CGT ACA TIC GIT ITA CAA TCG TCG GAA CTA CIT ITI TTA TCG CAA TCA GTA GAT TTA CCT GGT GGA ATG ACA GTT CTT GTA AGT AAA GAA AGT CAT CTA AAT GGA CCA CCT TAC TGT CAA GAA CAT TCA TTT CTT 160 150 170 AAA GAC AAA GAC GGT AAA TAC AGT CTA GAG GCA ACA GTA GAC AAG TIT CTG TIT CTG CCA TIT ATG TCA GAT CTC CGT TGT CAT CTG TTC 200 210 * * CTT GAG CTT AAA GGA ACT TCT GAT AAA AAC AAC GGT TCT GGA ACA GAA CTC GAA TTT CCT TGA AGA CTA TTT TTG TTG CCA AGA CCT TGT 240 CTT GAA GGT GAA AAA ACT GAC AAA AGT AAA GTA AAA TTA ACA ATT GAA CIT CCA CIT TIT TGA CIG TIT TCA TIT CAT TIT AAT TGT TAA 280 290 300 GCT GAT GAC CTA AGT CAA ACT AAA TIT GAA ATT TIC AAA GAA GAT CGA CTA CTG GAT TCA GTT TGA TTT AAA CTT TAA AAG TTT CTT CTA 330 340 350 * OCC AAA ACA TTA GTA TCA AAA AAA GTA ACC CTT AAA GAC AAG TCA COG TIT TGT AAT CAT AGT TIT TIT CAT TGG GAA TIT CTG TTC AGT 380 TCA ACA GAA GAA AAA TTC AAC GAA AAG GGT GAA ACA TCT GAA AAA AGT TGT CTT TTT AAG TTG CTT TTC CCA CTT TGT AGA CTT TTT 420 430 410 ACA ATA GTA AGA GCA AAT GGA ACC AGA CTT GAA TAC ACA GAC ATA TGT TAT CAT TCT CGT TTA CCT TGG TCT GAA CTT ATG TGT CTG TAT 470 480 AAA AGC GAT OGA TCC GGA AAA GCT AAA GAA GTT TTA AAA GAC TTT TIT TOG CTA CCT AGG CCT TIT CGA TIT CIT CAA AAT TIT CTG AAA 500 510 520 ACT CTT GAA GGA ACT CTA GCT GCT GAC GGC AAA ACA ACA TTG AAA TGA GAA CTT CCT TGA GAT CGA CGA CTG CCG TTT TGT TGT AAC TTT 570 560 GTT ACA GAA GGC ACT GTT GTT TTA AGC AAG AAC ATT TTA AAA TCC

K48/Tro OspA Wednesday, April 27, 1994 11:37 AM

CAA TGT CTT CCG TGA CAA CAA AAT TCG TTC TTG TAA AAT TTT AGG

590 600 610 620 630

GGA GAA ATA ACA GTT GCA CTT GAT GAC TCT GAC ACT ACT CAG GCT
CCT CTT TAT TGT CAA CGT GAA CTA CTG AGA CTG TGA TGA GTC CGA

. 640 650 660 670

ACT AAA AAA ACT OGA AAA TOG GAT TCA AAT ACT TCC ACT TTA ACA TGA TTT TTT TGA CCT TTT ACC CTA AGT TTA TGA AGG TGA AAT TGT

680 690 700 710 720

ATT AGT GTG AAT AGC AAA AAA ACT AAA AAC ATT GTA TIT ACA AAA TAA TCA CAC TTA TCG TIT TTT TGA TTT TTG TAA CAT AAA TGT TTT

730 740 750 760,

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GAA GAC ACA ATA ACA GTA CAA AAA TAC GAC TCA GCA GGC ACC AAT

CTT CTG TGT TAT TGT CAT GTT TTT ATG CTG AGT CGT CCG TGG TTA

770 780 790 800 810

CTA GAA GGC AAC GCA GTC GAA ATT AAA ACA CTT GAT GAA CTT AAA
GAT CTT CCG TTG CGT CAG CTT TAA TTT TGT GAA CTA CTT GAA TTT

820

AAC GCT TTA AAA TAG TTG CGA AAT TTT ATC

Figure 43 (2 of 2)

30 10 20 ATG AAA AAA TAT TTA TTG GGA ATA GGT CTA ATA TTA GCC TTA ATA TAC TIT TIT ATA AAT AAC CCT TAT CCA GAT TAT AAT CGG AAT TAT 60 GCA TGC AAG CAA AAT GTT AGC AGC CTT GAT GAA AAA AAC AGC GCT CGT ACG TTC GTT TTA CAA TCG TCG GAA CTA CTT TTT TTG TCG CGA 110 120. _____, 130 TCA GTA GAT TTG CCT GGT GAG ATG AAA GTT CTT GTA AGT AAA GAA AGT CAT CTA AAC GGA CCA CTC TAC TTT CAA GAA CAT TCA TTT CTT 140 150 160 170 AAA GAC AAA GAC OOT AAG TAC AOT CTA AAG OCA ACA GTA GAC AAG TIT CIG TIT CIG CCA TIC ATG TCA GAT TIC CGT TGT CAT CIG TIC 200 210 ATT GAG CTA AAA GGA ACT TCT GAT AAA GAC AAT GGT TCT GGA GTG TAA CTC GAT TIT CCT TGA AGA CTA TIT CTG TTA CCA AGA CCT CAC 240 CTT GAA GGT ACA AAA GAT GAC AAA AGT AAA GCA AAA TTA ACA ATT GAA CIT CCA TGT TIT CTA CIG TIT TCA TIT CGI TIT AAT TGT TAA 310 280 290 300 GCT GAC GAT CTA AGT AAA ACC ACA TTC GAA CTT TTA AAA GAA GAT CGA CTG CTA GAT TCA TTT TGG TGT AAG CTT GAA AAT TTT CTT CTA 350 340 330 GGC AAA ACA TTA GTG TCA AGA AAA GTA AGT TCT AGA GAC AAA ACA CCG TIT TGT AAT CAC AGT TCT TIT CAT TCA AGA TCT CTG TIT TGT 380 TCA ACA GAT GAA ATG TTC AAT GAA AAA GGT GAA TTG TCT GCA AAA AGT TGT CTA CTT TAC AAG TTA CTT TTT CCA CTT AAC AGA CGT TTT 420 430 440 450 ACC ATG ACA AGA GAA AAT GGA ACC AAA CTT GAA TAT ACA GAA ATG TGG TAC TGT TCT CTT TTA CCT TGG TTT GAA CTT ATA TGT CTT TAC 470 480 470 AAA AGC GAT GGA ACC GGA AAA GCT AAA GAA GTT TTA AAA AAG TTT TIT TOG CTA COT TGG COT TIT CGA TIT CIT CAA AAT TIT TIC AAA 530 510 520 500 ACT CTT GAA GGA AAA GTA GCT AAT GAT AAA GTA ACA TTG GAA GTA TGA GAA CTT CCT TTT CAT CGA TTA CTA TTT CAT TGT AAC CTT CAT 560 570 580 AAA GAA GGA ACC GTT ACT TTA AGT AAG GAA ATT GCA AAA TCT GGA . 20

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P-GAU/BO-OSPA Wednesday, April 27, 1994 11:22 AM

TIT CTT CCT TGG CAA TOA AAT TCA TTC CTT TAA CGT TTT AGA CCT

590 600 610 620 630

GAA GTA ACA GTT GCT CTT AAT GAC ACT AAC ACT ACT CAG GCT ACT CTT CAT TGT CAA CGA GAA TTA CTG TGA TTG TGA TGA GTC CGA TGA

640 650 660 670

680 690 700 710 720

AGT GTT AAC AGC AAA AAA ACT ACA CAA CTT GTG TTT ACT AAA CAA TCA CAA TTG TCG TTT TTT TGA TGT GTT GAA CAC AAA TGA TTT GTT

730 740 750 760

GAC ACA ATA ACT GTA CAA AAA TAC GAC TCC GCA GCT ACC AAT TTA CTG TGT TAT TGA CAT GTT TTT ATG CTG AGG CGT CCA TGG TTA AAT

770 780 790 800 810

GAA GGC ACA GCA GTC GAA ATT AAA ACA CTT GAT GAA CTT AAA AAC CTT CCG TGT CGT CAG CTT TAA TTT TGT GAA CTA CTT GAA TTT TTG

820

GCT TTA AAA TAG CGA AAT TTT ATC

Figure 44 (2 of 2)

10 ATG AAA AAA TAT TTA TTG GGA ATA GGT CTA ATA TTA GCC TTA ATA TAC TIT TIT ATA AAT AAC CCT TAT CCA GAT TAT AAT CGG AAT TAT 50 60 70 80 GCA TGT AAG CAA AAT GTT AGC AGC CTT GAC GAG AAA AAC AGC GTT CGT ACA TIC GIT TTA CAA TCG TCG GAA CTG CTC TTT TTG TCG CAA 110 120 -TCA GTA GAT TTG CCT GGT GAA ATG AAA GTT CTT GTA AGC AAA GAA AGT CAT CTA AAC GGA CCA CTT TAC TIT CAA GAA CAT TCG TIT CIT 170 180 ANA ANC ANA GAC GGC ANG TAC GAT CTA ATT GCA ACA GTA GAC. ANG TIT TIG TIT CIG CCG TIC ATG CTA GAT TAA CGT TGT CAT CTG TTC 200 210 CTT GAG CTT AAA GGA ACT TCT GAT AAA AAC AAT GGA TCT GGA GTA GAA CTC GAA TTT CCT TGA AGA CTA TTT TTG TTA CCT AGA CCT CAT 250 270 CTT GAA GCC GTA AAA GCT GAC AAA AGT AAA GTA AAA TTA ACA ATT GAA CIT CCG CAT TIT CGA CTG TIT TCA TIT CAT TIT AAT TGT TAA 280 290 300 TCT GAC GAT CTA GGT CAA ACC ACA CTT GAA GTT TTC AAA GAA GAT AGA CTG CTA GAT CCA GTT TGG TGT GAA CTT CAA AAG TTT CTT CTA 340 350 360 GGC AAA ACA CTA GTA TCA AAA AAA GTA ACT TCC AAA GAC AAG TCA CCG TTT TGT GAT CAT AGT TTT TTT CAT TGA AGG TTT CTG TTC AGT 390 400 TCA ACA GAA GAA AAA TTC AAT GAA AAA GGT GAA GTA TCT GAA AAA AGT TGT CTT CTT TTT AAG TTA CTT TTT CCA CTT CAT AGA CTT TTT 420 430 440 ATA ATA ACA AGA GCA AAT GGA ACC AAA CTT GAA TAT ACA GAA ATG TAT TAT TGT TCT CGT TTA CCT TGG TTT GAA CTT ATA TGT CTT TAC 460 470 480 . AAA AGC GAT GGA ACC GGA AAA GCT AAA GAA GTT TTA AAA AAG TTT TTT TCG CTA CCT TGG CCT TTT CGA TTT CTT CAA AAT TTT TTC AAA 500 510 520 530 ACT CTT GAA GGA AAA GTA GCT AAT GAT AAA GTA ACA TTG GAA GTA TGA GAA CTT CCT TTT CAT CGA TTA CTA TTT CAT TGT AAC CTT CAT 560 570 AAA GAA GGA ACC GIT ACT TTA AGT AAG GAA ATT TCA AAA TCT GGG

B31-PBK Mednesday, April 27, 1994 11:19 AK

TIT CTT CCT TGG CAA TGA AAT TCA TTC CTT TAA AGT TIT AGA CCC

590 600 610 620 630

GAA GIT TCA GTT GAA CTT AAT GAC ACT GAC AGT AGT GCT GCT ACT CTT CAA AGT CAA CTT GAA TTA CTG TGA CTG TCA TCA CGA CGA TGA

640 650 660 670

680 690 700 710 720

AGT GTG AAT AGC CAA AAA ACC AAA AAC CIT GTA TTC ACA AAA GAA TCA CAC TTA TCG GTT TTT TGG TTT TTG GAA CAT AAG TGT TTT CIT

730 740 750 760

GAC ACA ATA ACA GTA CAA AAA TAC GAC TCA GCA GGC ACC AAT CTA

770 780 790 800 810

GAA GGC AAA GCA GTC GAA ATT ACA ACA CTT AAA GAA CTT AAA AAC CTT CCG TTT CGT CAG CTT TAA TGT TGT GAA TTT CTT GAA TTT TTG

820

CT TTA AAA TAA CGA AAT TIT ATT

Figure 45 (2 of 2)

20 30 40 ATG AAA AAA TAT TTA TTG GGA ATA GGT CTA ATA TTA GCC TTA ATA TAC TIT TIT ATA AAT AAC CCT TAT CCA GAT TAT AAT CGG AAT TAT GCA TGC AAG CAA AAT GTT AGC AGC CTT GAT GAA AAA AAC AGC GCT CGT ACG TTC GTT TTA CAA TCG TCG GAA CTA CTT TTT TTG TCG CGA 120 TCA GTA GAT TTG CCT GGT GAG ATG AAA GTT CTT GTA AGT AAA GAA AGT CAT CTA AAC GGA CCA CTC TAC TTT CAA GAA CAT TCA TTT CTT 150 160 170 140 * AAA GAC AAA GAC GGT AAG TAC AGT CTA AAG GCA ACA GTA GAC AAG TIT CTG TIT CTG CCA TIC ATG TCA GAT TIC CGT TGT CAT CTG TIC 200 210 • . ATT GAG CTA AAA GGA ACT TCT GAT AAA GAC AAT GGT TCT GGA GTG TAA CTC GAT TTT CCT TGA AGA CTA TTT CTG TTA CCA AGA CCT CAC 230 240 250 260 270 CTT GAA GGT ACA AAA GAT GAC AAA AGT AAA GCA AAA TTA ACA ATT GAA CTT CCA TGT TTT CTA CTG TTT TCA TTT CGT TTT AAT TGT TAA 290 300 . GCT GAC GAT CTA AGT AAA ACC ACA TTC GAA CTT TTA AAA GAA GAT CGA CTG CTA GAT TCA TIT TGG TGT AAG CTT GAA AAT TIT CTT CTA 340 350 360 GGC AAA ACA TTA GTG TCA AGA AAA GTA AGT TCT AGA GAC AAA ACA CCG TTT TGT AAT CAC AGT TCT TTT CAT TCA AGA TCT CTG TTT TGT 390 380 400 TCA ACA GAT GAA ATG TTC AAT GAA AAA GGT GAA TTG TCT GCA AAA AGT TGT CTA CTT TAC AAG TTA CTT TTT CCA CTT AAC AGA CGT TTT 420 430 ACC ATG ACA AGA GAA AAT GGA ACC AAA CTT GAA TAT ACA GAA ATG TGG TAC TGT TCT CTT TTA CCT TGG TTT GAA CTT ATA TGT CTT TAC 480 AAA AGC GAT GGA ACC GGA AAA GCT AAA GAA GTT TTA AAA AAG TTT TIT TCG CTA CCT TGG CCT TTT CGA TTT CIT CAA AAT TIT TTC AAA 530 * * * 510 ACT CTT GAA GGA AAA GTA GCT AAT GAT AAA GTA ACA TTG GAA GTA TGA GAA CTT CCT TIT CAT CGA TTA CTA TTT CAT TGT AAC CTT CAT 56C 570 580 550 AAA GAA GGA ACC GTT ACT TTA AGT AAG GAA ATT TCA AAA TCT OGG

TIT CIT CCT TGG CAA TGA AAT TCA TTC CTT TAA AGT TTT AGA CCC 630 610 • GAA GTT TCA GTT GAA CTT AAT GAC ACT GAC AGT AGT GCT GCT ACT CTT CAA AGT CAA CTT GAA TTA CTG TGA CTG TCA TCA CGA CGA TGA 650 * 660 4 AAA AAA ACT GCA GCT TGG AAT TCA AAA ACT TCC ACT TTA ACA ATT TIT TIT TGA CGT CGA ACC TTA AGT TTT TGA AGG TGA AAT TGF TAA 710 690 680 AGT GTG AAT AGC CAA AAA ACC AAA AAC CTT GTA TTC ACA AAA GAA TCA CAC TTA TCG CTT TTT TCG TTT TTG GAA CAT AAG TGT TTT CTT 760 750 740 • GAC ACA ATA ACA GTA CAA AAA TAC GAC TCA GCA OGC ACC AAT CTA CTG TGT TAT TGT CAT GTT TTT ATG CTG AGT CGT CCG TGG TTA GAT 790 780 770 ŧ GAA GGC AAA GCA GTC GAA ATT ACA ACA CTT AAA GAA CTT AAA AAC CTT CCG TTT CGT CAG CTT TAA TGT TGT GAA TTT CTT GAA TTT TTC 820 OCT TTA AAA TAA CGA AAT TTT ATT

A. CLASSIFICATION OF SY IPC 6 C12N15/3 C07K16/12

T MATTER C12N15/62

C07K14/20

A61K3

G01N33/50

According to International Patent Classification (IPC) or to both national classification and IPC

#### **B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols) IPC 6 CO7K C12N A61K G01N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

C. DOCUI	MENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	MOLECULAR MICROBIOLOGY, vol.6, no.20, 1992 pages 3031 - 3040 ROSA P. A. ET AL. 'Recombination between genes encoding major outer surface proteins A and B of Borrelia burgdorferi' see the whole document	1,6,7,9, 22,27, 28,30,45

X Further documents are listed in the continuation of box C.	X Patent family members are listed in annex.
*Special categories of cited documents:  A* document defining the general state of the art which is not considered to be of particular relevance  E* earlier document but published on or after the international filing date  L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)  O* document referring to an oral disclosure, use, exhibition or other means  P* document published prior to the international filing date but later than the priority date claimed	T later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention  'X' document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone  'Y' document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.  '&' document member of the same patent family
Date of the actual completion of the international search	Date of mailing of the international search report
24 February 1995	03. 03. 95
Name and mailing address of the ISA  European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Espen, J

Form PCT/ISA/210 (second sheet) (July 1992)



GINSBERG H. S. ET AL. (EDS.) 'Vaccines 93.  Modern approaches to new vaccines including prevention of AIDS. Tenth annual meeting, Cold Spring Harbor, New York, USA, September 1992.'  1993, COLD SPRING HARBOR LABORATORY PRESS, NEW YORK cited in the application McGrath B. C. et al.: 'Biochemical and biophysical characterization of the major outer surface protein from north american and european isolates of Borrelia burgdorferi' see page 365 - page 370 see page 369, last paragraph  X JOURNAL OF BACTERIOLOGY, vol.175, no.9, May 1993 pages 2516 - 2522 KITTEN T. ET AL. 'Intragenic recombination and a chimeric outer membrane protein in the relapsing fever agent Borrelia hermsii' see the whole document	Gluson of document.  D.X  GINSBERG H. S. ET AL. (EDS.) 'Vaccines 93.  Modern approaches to new vaccines including prevention of AIDS. Tenth annual meeting, Cold Spring Harbor, New York, USA, September 1992.'  1993, COLD SPRING HARBOR LABORATORY PRESS, NEW YORK cited in the application McGrath B. C. et al.: 'Biochemical and biophysical characterization of the major outer surface protein from north american and european isolates of Borrelia burgdorferi' see page 365, last paragraph  X  JOURNAL OF BACTERIOLOGY, vol.175, no.9, May 1993 pages 2516 - 2522  KITTEN T. ET AL. 'Intragenic recombination and a chimeric outer membrane protein in the relapsing fever agent Borrelia hermsii' see the whole document  Y  WO,A,93 08306 (SYMBICOM AKTIEBOLAG) 29  April 1993  see claims 44-46  Y  WO,A,91 13630 (THE UNITED STATES OF AMERICA) 19 September 1991  see page 8; figures 11,13  P,X  WO,A,94 20536 (THE REGENTS OF THE UNIVERSITY OF CALIFORNIA) 15 September 1994	Gluson of document.  D.X  GINSBERG H. S. ET AL. (EDS.) 'Vaccines 93.  Modern approaches to new vaccines including prevention of AIDS. Tenth annual meeting, Cold Spring Harbor, New York, USA, September 1992.'  1993, COLD SPRING HARBOR LABORATORY PRESS, NEW YORK cited in the application McGrath B. C. et al.: 'Biochemical and biophysical characterization of the major outer surface protein from north american and european isolates of Borrelia burgdorferi' see page 369, last paragraph  X  JOURNAL OF BACTERIOLOGY, vol.175, no.9, May 1993 pages 2516 - 2522  KITTEN T. ET AL. 'Intragenic recombination and a chimeric outer membrane protein in the relapsing fever agent Borrelia hermsii' see the whole document  Y  WO,A,93 08306 (SYMBICOM AKTIEBOLAG) 29  April 1993  see claims 44-46  Y  WO,A,91 13630 (THE UNITED STATES OF AMERICA) 19 September 1991  see page 8; figures 11,13  P,X  WO,A,94 20536 (THE REGENTS OF THE UNIVERSITY OF CALIFORNIA) 15 September 1994			PC1703 3471	
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